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Conservation of inner nuclear membrane targeting sequences in mammalian Pom121 and yeast Heh2 membrane proteins

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ABSTRACT Endoplasmic reticulum–synthesized membrane proteins traffic through the nuclear pore complex (NPC) en route to the inner nuclear membrane (INM). Although many membrane proteins pass the NPC by simple diffusion, two yeast proteins, ScSrc1/ScHeh1 and ScHeh2, are actively imported. In these proteins, a nuclear localization signal (NLS) and an intrinsically disordered linker encode the sorting signal for recruiting the transport factors for FG-Nup and RanGTP-dependent transport through the NPC. Here we address whether a similar import mechanism applies in metazoans. We show that the (putative) NLSs of metazoan HsSun2, MmLem2, HsLBR, and HsLap2 β are not sufficient to drive nuclear accumulation of a membrane protein in yeast, but the NLS from RnPom121 is. This NLS of Pom121 adapts a similar fold as the NLS of Heh2 when transport factor bound and rescues the subcellular localization and synthetic sickness of Heh2 Δ NLS mutants. Consistent with the conservation of these NLSs, the NLS and linker of Heh2 support INM localization in HEK293T cells. The conserved features of the NLSs of ScHeh1, ScHeh2, and RnPom121 and the effective sorting of Heh2-derived reporters in human cells suggest that active import is conserved but confined to a small subset of INM proteins.

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L.M.V. and A.K. initiated the project and wrote the manuscript together with all of the other authors. A.K. performed all experiments except the biochemical and structural studies in Figure 2, which were performed and analyzed by R.K.L. and G.C. The experiments in HEK293T cells in Figure 6, B and C, and Supplemental Figure S2 were designed by A.K., M.F., and N.J.B., performed by A.K. and N.J.B., and analyzed by A.K. and M.F. Experiments in Figures 5 and 6A were performed with the help of V.v.d.B.; A.S. performed experiments in Figure 4.

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Abbreviations used: DAM, DNA adenine methyltransferase; ER, endoplasmic reticulum; FKBP, FK506-binding protein; FL, full length; FRB, FKBP12-rapamycin binding domain; GFP, green fluorescent protein; IBB, importin β binding; INM, inner nuclear membrane; NE, nuclear envelope; NLS, nuclear localization signal; NPC, nuclear pore complex; Nup, nucleoporin; ONM, outer nuclear membrane.

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INTRODUCTION

The gateway between the cytoplasm and the nuclear interior is formed by the nuclear envelope (NE)–embedded nuclear pore complex (NPC) through which bidirectional transport between the two compartments occurs. Small solutes and proteins readily diffuse through the NPC, whereas diffusion of larger macromolecular complexes is slow or prevented. The presence of a nuclear localization signal (NLS) or nuclear export signal recognized by transport receptors permits transport using an energy-dependent mechanism (Chook and Suel, 2011) and also allows the passage of larger molecules. Specific interactions of these transport receptors with phenylalanine-glycine repeats of nucleoporins (Nups) filling the central channel of the NPC mediate the transport of soluble cargoes (Fiserova et al., 2010; Peleg and Lim, 2010; Yang, 2013), and receptor–cargo binding and release are dictated by a gradient of RanGTP across the NPC (Kalab et al., 2002; Fried and Kutay, 2003; Madrid and Weis, 2006; Cook et al., 2007).

In postmitotic cells and cells undergoing a closed mitosis, inner nuclear membrane (INM)-localized integral membrane proteins also have to find their way through the NPC. Upon synthesis, polytopic membrane proteins are first incorporated into the endoplasmic reticulum (ER) membrane and travel via the interconnected outer nuclear membrane (ONM) and pore membrane (which lines the NPC) to the INM. The INM contains a unique set of integral membrane proteins (Schirmer *et al.*, 2003). In contrast to the extensively studied energy-dependent import mechanisms for soluble proteins, little is known about active transport of INM membrane proteins.

At first, accumulation of INM proteins in metazoans was explained by diffusion and retention (Powell and Burke, 1990; Ellenberg *et al.*, 1997). Here membrane proteins diffuse between the ONM and the INM, while their extralumenal domain slides through the lateral channel of the NPC, and are retained (and accumulated) at the INM by interaction with chromatin or nuclear proteins, most notably lamins (Soullam and Worman, 1993; Furukawa *et al.*, 1998; Lin *et al.*, 2005). This mode of import is limited to membrane proteins with an extralumenal domain small enough to fit the lateral channel, which has a width of ~10 nm (Hinshaw *et al.*, 1992; Ohba *et al.*, 2004; Bui *et al.*, 2013). Later it was shown that metabolic energy is required for the translocation of a group of integral membrane proteins to the INM (Ohba *et al.*, 2004). The dynamics of 15 NE transmembrane proteins in the NE-ER network showed that ATP- and Ran-dependent translocation mechanisms are distinct and not used by all inner nuclear membrane proteins (Zuleger *et al.*, 2011).

In *Saccharomyces cerevisiae*, Heh2 and Src1/Heh1 are actively transported to the INM by karyopherins 60 and 95 (Kap60/Kap95), the yeast homologues of importin α and importin β , respectively (King *et al.*, 2006). The NLSs of Heh1 and Heh2 are separated from the transmembrane domain by a region that is intrinsically disordered (ID; Meinema *et al.*, 2011). This ID linker region and an NLS are required and sufficient to accumulate a membrane protein at the INM in the absence of retention, in that a reporter protein consisting of just these domains is mobile in the INM (Meinema *et al.*, 2011).

Importin α consists of a tandem array of 10 Armadillo (ARM) repeats that together form a superhelical structure with a major NLS binding pocket between ARM repeats 2–4 and a minor NLS binding pocket between ARM repeats 7 and 8 (Conti *et al.*, 1998). In both binding sites, five contact points with NLSs are identified, P1–P5 at the major binding site and P1'–P5' at the minor binding pocket (Conti and Kuriyan, 2000; Fontes *et al.*, 2003; Giesecke and Stewart, 2010; Marfori *et al.*, 2012; Roman *et al.*, 2013). An N-terminal auto-inhibitory domain of importin α , called the importin β -binding domain (IBB domain; reviewed in Lott and Cingolani, 2011), and the NLSs compete for the same binding pocket on importin α . Once importin β binds to the IBB domain, the binding sites on importin α are exposed and able to interact with an NLS (Gorlich *et al.*, 1996; Moroiaru *et al.*, 1996; Lott *et al.*, 2010). The NLS of Heh2 (residues 102–131) is distinctive from other NLSs in that it binds to full-length importin α in the absence of importin β (King *et al.*, 2006). Further, the Heh1 and Heh2 NLS binds importin α in the absence of importin β , and both adopt an IBB-like fold while bound to Kap60 (Lokareddy *et al.*, 2015).

NLS sequences are predicted or described in many metazoan INM proteins, but whether a similar metabolic energy- and Ran and Kap-dependent import mechanism as described in yeast applies is not known. Indeed, these (potential) importin α/β interacting sequences could also act in nuclear envelope reformation, in which importin α and importin β act in mitotic spindle formation

and NE assembly. Here the Ran guanine exchange factor regulator of chromosome condensation (RCC1) generates GTP-bound Ran GTPase (RanGTP) and is chromatin associated, resulting in high concentrations of RanGTP locally around chromosomes (Kalab *et al.*, 2002; Zhang *et al.*, 2002a; Hutchins *et al.*, 2009). RanGTP induces the dissociation of the importin α/β from NLS-containing proteins that function as spindle assembly factors (e.g., NUMA, TPX2, and the kinesin XCTK2), which are inactive when bound to importin α (Gruss *et al.*, 2001; Schatz *et al.*, 2003; Ems-McClung *et al.*, 2004). How importin β contributes to the reformation of the NE is not completely elucidated, but in vitro studies with cell-free systems made from *Xenopus laevis* egg extracts show that it might be involved in recruitment of FxFG-containing Nups (Zhang *et al.*, 2002b) and lamin-B receptor (LBR)-containing membrane vesicles to the decondensing chromatin (Ma *et al.*, 2007). Other integral membrane proteins contribute to the reformation of the NE as well, via direct interaction with DNA (Ulbert *et al.*, 2006), or by recruitment mediated by importin α/β .

Here we studied five INM proteins that contain putative NLSs and ID linker domains: Sun2, Lem2, LBR, Lap2 β , and Pom121. We tested the ability of these putative NLSs to target a membrane protein to the inner membrane in yeast and found that only the NLS of Pom121 does. Structural, biochemical, and in vivo studies reveal that the first two boxes of positive residues (residues 290–320) in the NLS region of Pom121 (residues 290–484) share with Heh1 and Heh2 an IBB-like fold when bound to importin α . Pom121NLS₂₉₀₋₃₂₆ is able to restore the INM localization of Heh2 Δ h2NLS and rescues cellular sickness of Heh2 Δ h2NLS in a strain lacking Nup84 as well. Consistent with the evolutionary conservation of these NLSs, we show that the NLS of Heh2 supports INM localization of a membrane protein in HEK293T cells. This suggests that active import of INM proteins may be conserved but confined to a small subset of the inner nuclear membrane proteins.

RESULTS

Pom121NLS mediates INM import of a membrane reporter protein in yeast

In *S. cerevisiae*, it was shown for the membrane protein Heh2 that import to the INM depends on the presence of an NLS (King *et al.*, 2006) that adopts an IBB-like fold (Lokareddy *et al.*, 2015) and on an intrinsically disordered linker region (L) that creates distance between the NLS and the transmembrane (TM) segment (Meinema *et al.*, 2011); this domain composition is collectively called NLS-L-TM. Several metazoan INM proteins with described or putative NLSs also encode regions that are predicted to be intrinsically disordered, based on predictions from FoldIndex (Prilusky *et al.*, 2005). We found a putative NLS-L-TM signature in *Homo sapiens* Sun2, LBR, and Lap2 β and *Mus musculus* Lem2, and we studied whether the NLSs in these domains are sufficient to support active INM import in baker's yeast. The choice of baker's yeast is related to its closed mitosis, meaning that the NE stays intact during anaphase, and in this way the sole route to the INM is via the NPCs. In metazoan systems, it is more difficult to distinguish INM import via the NPC from other (NLS-mediated) targeting mechanisms that may occur when the NE is disintegrated.

We constructed reporter proteins in which the mammalian NLSs were fused to green fluorescent protein (GFP) and to the ID linker region and first TM segment of Heh2 (Figure 1A, GFP-NLS-L-TM). The localization of the membrane reporter proteins was visualized with fluorescence microscopy, and all four reporters with NLSs derived from Sun2, Lem2, LBR, and Lap2 β localized to the NE-ER network as the TM segment alone does (Figure 1A), whereas the

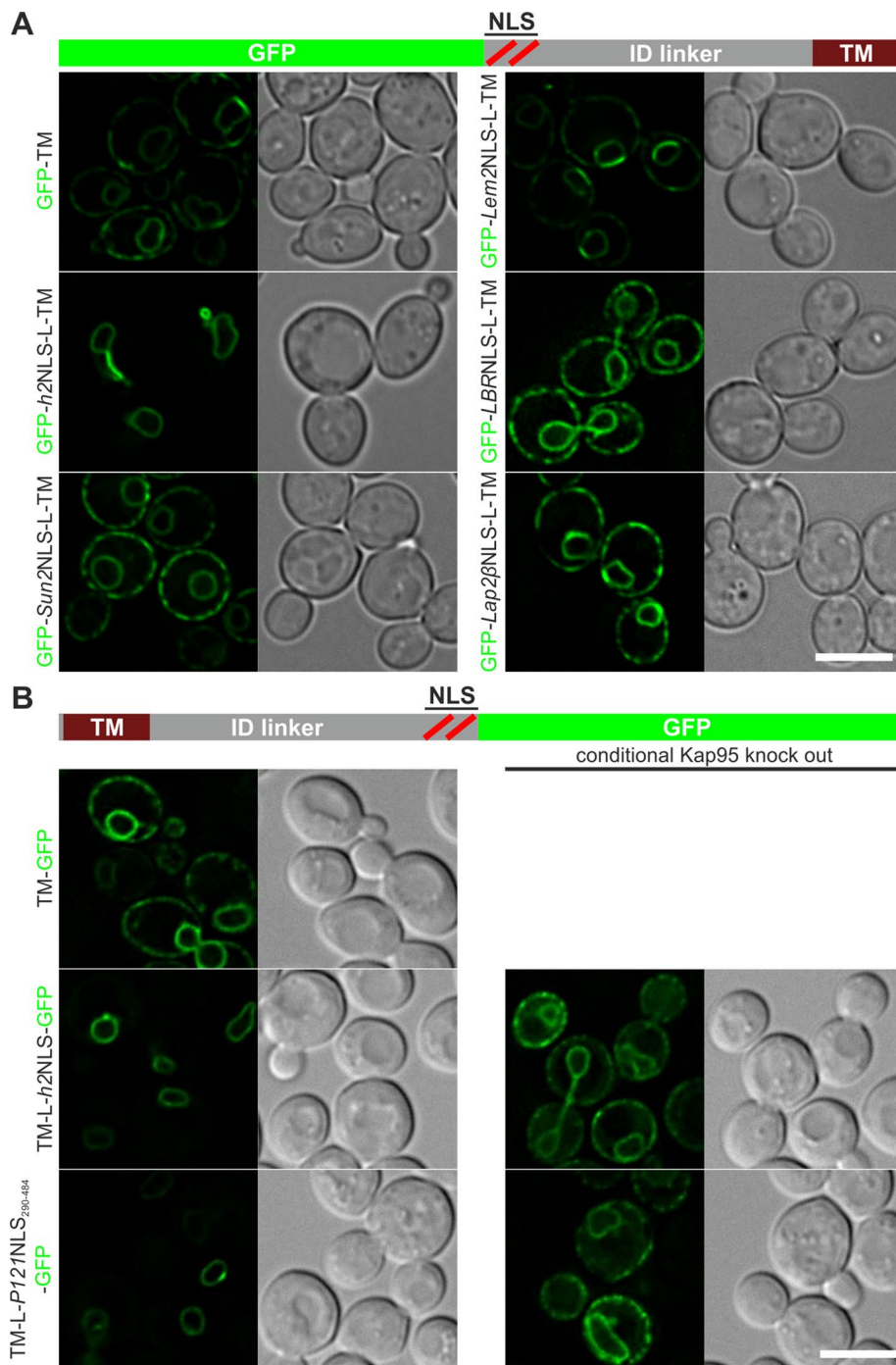


FIGURE 1: Localization of membrane-embedded reporter proteins encoding mammalian NLS sequences. (A) N-terminal GFP fusions of (putative) NLS regions of Sun2, Lem2, LBR, or Lap2 β , the intrinsically disordered linker region (L), and first transmembrane segment (TM) of Heh2 localize to the NE-ER network, like the TM of Heh2 (GFP-TM), and do not mediate nuclear accumulation like the NLS of Heh2 (GFP-h2NLS-L-TM). (B) Whereas a C-terminally GFP fusion of a TM localizes to the NE-ER network (TM-GFP), fusion of this TM to the ID linker region of Heh2 and the NLS regions of Heh2 or Pom121 (TM-L-h2NLS-GFP and TM-L-P121NLS₂₉₀₋₄₈₄-GFP, respectively) results in strong NE accumulation of the protein. This accumulation is lost in the conditional Kap95 knockout. Scale bars, 5 μ m.

same reporter with the Heh2 NLS (h2NLS) localized to the nuclear envelope. We also fused the complete predicted NLS-ID linker region of Sun2, Lem2, LBR, and Lap2 β to the TM segment of Heh2, but none of these regions was able to mediate INM import in yeast (un-

published data). This does not negate a contribution of the NLSs to targeting in vivo. Instead, it shows that these mammalian NLSs are not a functional equivalent of the h2NLS, as they are insufficient to support active INM import, leading to a steady-state accumulation of the membrane protein at the INM.

Another candidate with an established NLS region that we tested was Pom121 (Doucet *et al.*, 2010; Yavuz *et al.*, 2010; Funakoshi *et al.*, 2011). Pom121 is a single-pass membrane protein with a small luminal domain at the N-terminus and a large C-terminal extraluminal domain. The NLS region of Pom121, containing four or five boxes of basic residues, is highly conserved among species and encodes at least two bipartite NLSs (Yavuz *et al.*, 2010; Funakoshi *et al.*, 2011). The region between the TM domain and the NLS region is predicted to be at least in part disordered but also includes a more hydrophobic domain involved in targeting (Funakoshi *et al.*, 2011). We did not study this region but instead focused on the NLS. In Pom121, the topology is predicted as TM-L-NLS (Soderqvist and Hallberg, 1994), rather than NLS-L-TM as in Heh1 and Heh2. It was unknown whether reporters with this topology could support import at all, and this was tested first. We show that whereas a control reporter (TM-GFP) localized to the NE-ER network, a reporter with the ID linker and NLS of Heh2 (TM-L-h2NLS-GFP) accumulated at the INM (Figure 1B). To prove that this localization was Kap95 dependent, we studied its localization in a conditional Kap95-knockout strain. In this strain, Kap95 was tagged with FKBP12-rapamycin binding domain (FRB), which forms a stable heterodimer with Pma1-fused FK506-binding protein (FKBP) in the presence of rapamycin, resulting in trapping of Kap95 at the plasma membrane (Haruki *et al.*, 2008) and abolishing Kap60/Kap95-mediated transport over the NPC (Meinema *et al.*, 2011). Indeed, INM localization of TM-L-h2NLS-GFP depends on the presence of Kap95, as addition of rapamycin resulted in the appearance of ER-localized TM-L-h2NLS-GFP (Figure 1B). Consistent with previous findings that the ID linker length between the NLS and the transmembrane segment scales with the level of accumulation at the INM (Meinema *et al.*, 2011), we observed that accumulation is abolished when the length of the ID linker region in TM-L-h2NLS-GFP is reduced to 53 residues (Supplemental Figure S1A; TM-L53-h2NLS-GFP). We conclude that reporters with N- or C-terminal h2NLS and ID linker sorting signals are imported to the INM of yeast in a Kap60/Kap95-dependent manner.

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Next, to investigate whether the NLS of *Rattus norvegicus* (Rn) Pom121 supported INM import, we replaced the NLS of Heh2 for the NLS region of Rn Pom121 (residues 290–484), including multiple potential NLSs (Figure 1B; TM-L-Pom121NLS₂₉₀₋₄₈₄-GFP), and determined its subcellular localization. Clearly, the presence of this NLS region also resulted in Kap95-mediated INM localization of the TM reporter protein. Of the mammalian NLSs tested so far, only the NLS of Pom121 was able to mediate Kap60/Kap95-dependent INM targeting of a membrane protein in yeast.

Pom121NLS resembles the membrane protein NLS of Heh2

To visualize the interaction of Rn Pom121NLS with importin α 1, we focused on the NLS regions encoded by residues 291–320. We coexpressed in bacteria plasmids encoding Δ IBB-importin α 1 and GST-tagged Pom121NLS (residues 291–320), and captured a stoichiometric complex of the two proteins on glutathione beads. Coexpression prevented proteolytic degradation of Pom121NLS₂₉₁₋₃₂₀ and was instrumental in purifying milligram quantity of homogeneous complex that we used to grow high-quality crystals. The structure of Δ IBB-importin α 1 bound to Pom121NLS₂₉₁₋₃₂₀ was solved by molecular replacement and refined to an $R_{\text{work}}/R_{\text{free}}$ of 18.2/20.9% at 1.8-Å resolution (Supplemental Table S1). Pom121NLS₂₉₁₋₃₂₀ adopts an S-shaped conformation that binds the concave surface of importin α 1 Arm core, burying 2960 Å² of solvent-accessible surface area (Figure 2A). Twenty-two Pom121NLS₂₉₁₋₃₂₀ residues make close contact with as many as 59 residues in the Arm core, which are mainly clustered at the major (Arm 2–4) and minor (Arm 7 and 8) NLS-binding pockets and with sporadic interactions between Arm 5 and 6. Of note, Pom121NLS's first basic box makes four strong contacts at the minor NLS pocket, where the basic stretch 294-KKKR-297 occupies positions P1'–P4', as seen for membrane protein NLS of Heh1 (174-KKRR-177) and Heh2 (102-KRKR-105; Lokareddy et al., 2015). Similarly, at the major NLS-binding pocket, Pom121NLS₂₉₁₋₃₂₀ inserts four basic side chains at sites P2–P5 (313-KRRR-316) and makes additional contacts at P1 and P6 with N312 and H317, respectively. The average *B* factor of Pom121NLS₂₉₁₋₃₂₀ boxes interacting at minor (39.8 Å²) and major (19.7 Å²) NLS-binding pockets is comparable to that of importin α Arm core (28.6 Å²), consistent with the high avidity for this bipartite NLS for importin α . In contrast, the 14-residue spacer between NLS boxes (298-TVAEDQLHLDGQE-311) has significantly weaker electron density (visible continuously only by blurring the *B* factor) and adopts a random coiled conformation. Overall Pom121NLS₂₉₁₋₃₂₀ binding to importin α Arm core is stabilized by 45 hydrogen bonds, seven salt bridges, and a handful of hydrophobic and cation π (Korner et al., 2003) contacts involving importin α -conserved tryptophans.

Structural alignment of Pom121NLS₂₉₁₋₃₂₀ with NLSs visualized crystallographically in complex with importin α /Kap60 (Supplemental Table S2) revealed the position of critical residues at P2 and P2'. Pom121NLS₂₉₁₋₃₂₀ inserts a lysine at position P2 (K313) in the major NLS-binding pocket, as observed for the vast majority of classical and nonclassical NLSs (Kalderon et al., 1984). In contrast, unlike most NLSs, which make strong contacts at the minor NLS box (Kosugi et al., 2009; Giesecke and Stewart, 2010; Lott et al., 2011; Chang et al., 2013; Pang and Zhou, 2014; Pumroy et al., 2015), Pom121NLS₂₉₁₋₃₂₀ inserts a lysine at P2' (K295) as opposed to an arginine. The structural alignment also showed that Pom121NLS₂₉₁₋₃₂₀ shares striking structural resemblance to the IBB domain of importin α in its inhibitory conformation and to the Heh2 NLS (h2NLS). Pom121NLS, IBB, and h2NLS have superimposable traces (Figure 2B) and diverge only in the variable region between NLS boxes (residues 299–311 in Pom121NLS₂₉₁₋₃₂₀), which makes

minimal contacts with the Arm core. As observed for h2NLS, Pom121NLS₂₉₁₋₃₂₀ does not associate directly with importin β (unpublished data), suggesting that this NLS cannot adopt the helical fold essential for IBB association with importin α (Lott et al., 2010).

The striking similarity of Pom121NLS₂₉₁₋₃₂₀ to the IBB domain prompted us to determine whether Pom121NLS can bypass IBB autoinhibition, as found for h2NLS and, to a lesser extent, h1NLS (King et al., 2006; Lokareddy et al., 2015). To test this hypothesis, we used an on-bead binding assay in which glutathione *S*-transferase (GST)-tagged full-length (FL)-importin α 1 and GST- Δ IBB-importin α 1 were immobilized on glutathione beads and incubated with a twofold molar excess of maltose binding protein (MBP)-tagged Pom121NLS₂₉₁₋₃₂₀, NPNLS (a negative control for IBB displacement), or h2NLS (a positive control for IBB displacement). Of note, MBP-Pom121NLS₂₉₁₋₃₂₀ was minimally able to overcome autoinhibition, and <20% of the starting material was recovered bound to FL-importin α 1 beads after 15 min of incubation (Figure 2C), comparable to the negative control NPNLS, which is autoinhibited by the IBB domain (Lokareddy et al., 2015; Pumroy et al., 2015). In contrast, as previously shown, h2NLS completely bypassed IBB autoinhibition, binding importin α 1 Arm core equally in the presence or absence of IBB (Figure 2C). Mutation at P2' and P2 completely disrupted the interaction of Pom121NLS₂₉₁₋₃₂₀ with FL-importin α 1, indistinguishable from a double mutant at P2'/P2 (Figure 2D). A single mutation at P2 was also sufficient to disrupt Pom121NLS₂₉₁₋₃₂₀ binding to Δ IBB-importin α 1, suggesting the interaction of this NLS with the Arm-core is cemented by the major NLS-binding box, as observed in classical NLS (Figure 2D). Thus Pom121NLS₂₉₁₋₃₂₀ adopts an IBB-like structure that combines binding determinants seen in the recognition of classical NLSs and a deeper interaction at the minor NLS-binding site as observed for h2NLS.

Pom121NLS₂₉₀₋₃₂₆ is sufficient to mediate INM import of a membrane-embedded protein, and interactions at P2 and P2' positions are critical

To further study the characteristics of the IBB-like region of Pom121NLS, we created a reporter protein that was fused to only the IBB-like region of Pom121NLS (TM-L-P121NLS₂₉₀₋₃₂₆-GFP; Figure 3A). TM-L-Pom121NLS₂₉₀₋₃₂₆-GFP accumulated at the INM, although a fraction of the protein was observed in the ER. As expected, the INM accumulation was abolished upon depletion of Kap95. The introduction of an alanine at P2 or P2' (Figure 3B; TM-L-P121NLS₂₉₀₋₃₂₆ P2-GFP and P2'-GFP, respectively) disrupted the INM targeting of the reporter protein. This shows that interaction of the P2 position, as well as of the P2' position, of Pom121NLS₂₉₀₋₃₂₆ with importin α is required to mediate efficient INM import of a membrane-embedded protein, consistent with the structural data. Here the IBB-like NLS in the region of Pom121NLS (residues 291–320) differs from what was observed for h2NLS, for which binding of the P2' position contributed more significantly to INM import of the reporter membrane protein (Lokareddy et al., 2015).

The Pom121NLS region encodes multiple NLSs (Doucet et al., 2010; Yavuz et al., 2010; Funakoshi et al., 2011). Indeed, the substitutions at P2 or P2' (unpublished data), as well as the combination of these (Figure 3C; TM-L-P121NLS₂₉₀₋₄₈₄ P2/P2'-GFP), left the Kap95-dependent INM localization of the protein unaltered. In addition, removal of the IBB-like NLS of the NLS region of Pom121 (Figure 3C; TM-L-P121NLS₃₂₃₋₄₈₄-GFP) did not influence the Kap95-dependent import of the reporter protein. Clearly, besides Pom121NLS₂₉₁₋₃₂₀, the Pom121NLS region contains at least one alternative importin-binding site that is recognized in yeast and mediates INM import.

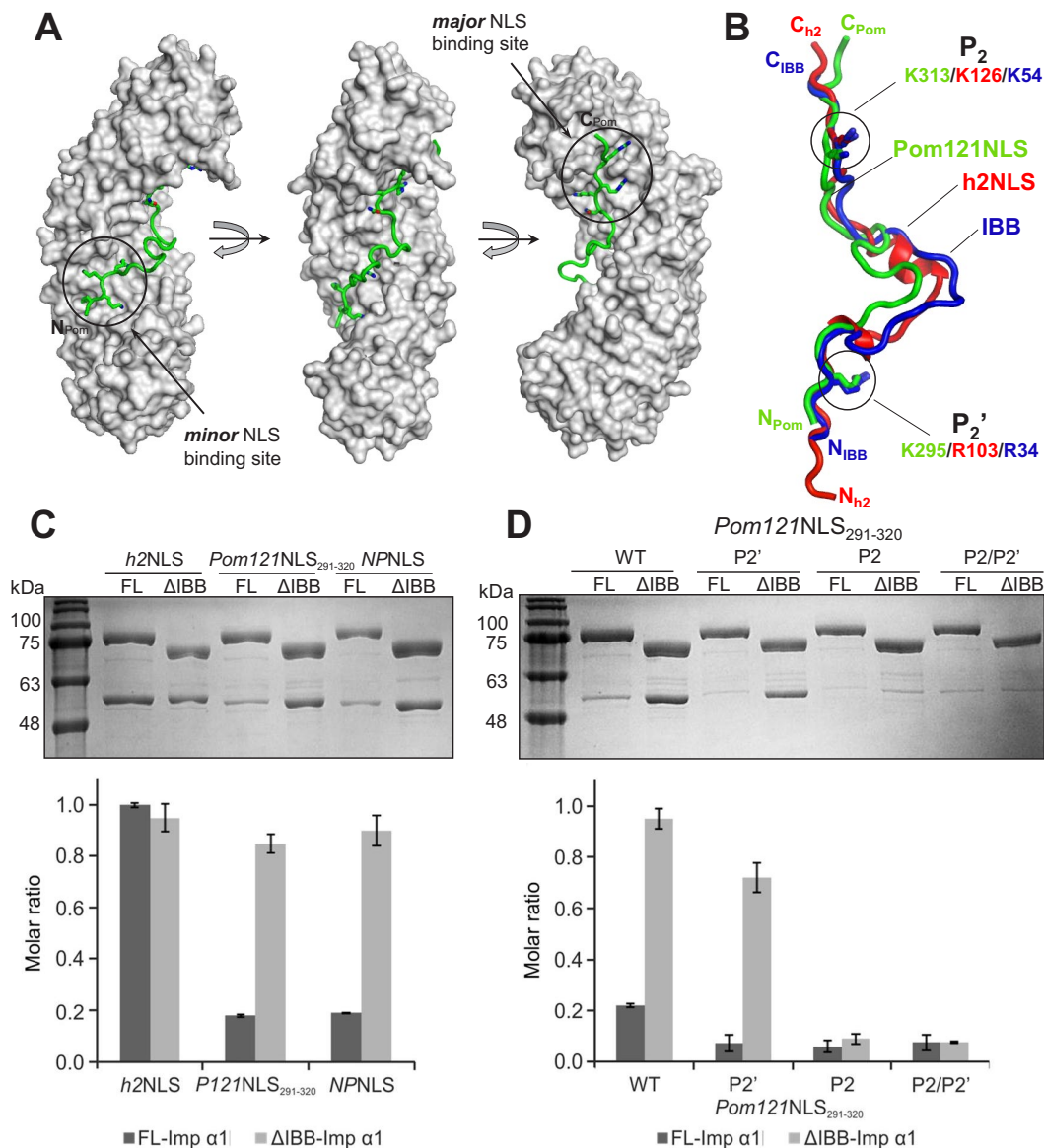


FIGURE 2: Structural and biochemical analysis of *Pom121NLS* bound to importin α 1. (A) Crystal structure of Δ IBB-importin α 1 (gray surface) in complex with *Pom121NLS*₂₉₁₋₃₂₀ (green) shown in three views rotated clockwise by 90°. (B) Superimposition of Δ IBB-importin α 1 bound to *Pom121NLS*₂₉₁₋₃₂₀ with Δ IBB-Kap60 bound to the *h2NLS* (Lokareddy et al., 2015) or as FL-Kap60 (Protein Data Bank ID 1WA5). *Pom121NLS*₂₉₁₋₃₂₀ and *h2NLS* are green and red, respectively, and the IBB domain is blue. For clarity, importin α 1 and Kap60 are omitted, and only the NLSs are shown. Residues at *P2'* and *P2* are shown as sticks. (C) Pull-down analysis and quantification of the interaction of GST-tagged importin α 1 lacking the IBB (Δ IBB) or FL immobilized on glutathione beads and incubated with MBP-tagged *h2NLS*, *Pom121NLS*₂₉₁₋₃₂₀, and *NP-NLS*. (D) Pull-down analysis and quantification of the interaction of GST- Δ IBB-importin α 1 or FL-importin α 1 with WT *Pom121NLS*₂₉₁₋₃₂₀ and mutants at *P2'*, *P2*, and *P2/P2'*. Pull downs in C and D are shown as mean \pm SD for three experiments.

Next we looked for evidence of similarity between the *h2NLS* and *Pom121NLS*₂₉₀₋₃₂₆ in the context of an actual inner membrane protein. First, we addressed whether replacement of the *h2NLS* in the endogenous *Heh2* gene for *Pom121NLS*₂₉₀₋₃₂₆ affects the subcellular localization. GFP-*Heh2* Δ *h2NLS* expressed from its endogenous promoter is mislocalized to the peripheral ER (Figure 4A; GFP-*Heh2* Δ *h2NLS*). Introducing *Pom121NLS*₂₉₀₋₃₂₆ rescues the localization at the NE: the localization of GFP-*Heh2*(Δ *h2NLS*, *Pom121NLS*₂₉₀₋₃₂₆) is indistinguishable from that of GFP-*Heh2* (Figure 4A). We also assessed synthetic lethality/sickness of yeast strains that combine mutations of NPC

components with mutations in the NLS region of *Heh2*, based on the previous finding that a double mutant lacking *Heh2* and *Nup84* is not viable (Yewdell et al., 2011). Strains lacking *Nup84* and expressing GFP-*Heh2*, GFP-*Heh2*(Δ *h2NLS*, *Pom121NLS*₂₉₀₋₃₂₆), or GFP-*Heh2*(Δ *h2NLS*) were tested, and whereas the double mutant expressing GFP-*Heh2*(Δ *h2NLS*) was synthetic sick, the double mutants expressing GFP-*Heh2* or GFP-*Heh2*(Δ *h2NLS*, *Pom121NLS*₂₉₀₋₃₂₆) grew well (Figure 4B). Overall these studies show the importance and similarity of the *Pom121NLS*₂₉₀₋₃₂₆ and the NLS of *Heh2* in vivo and in the context of full-length *Heh2*.

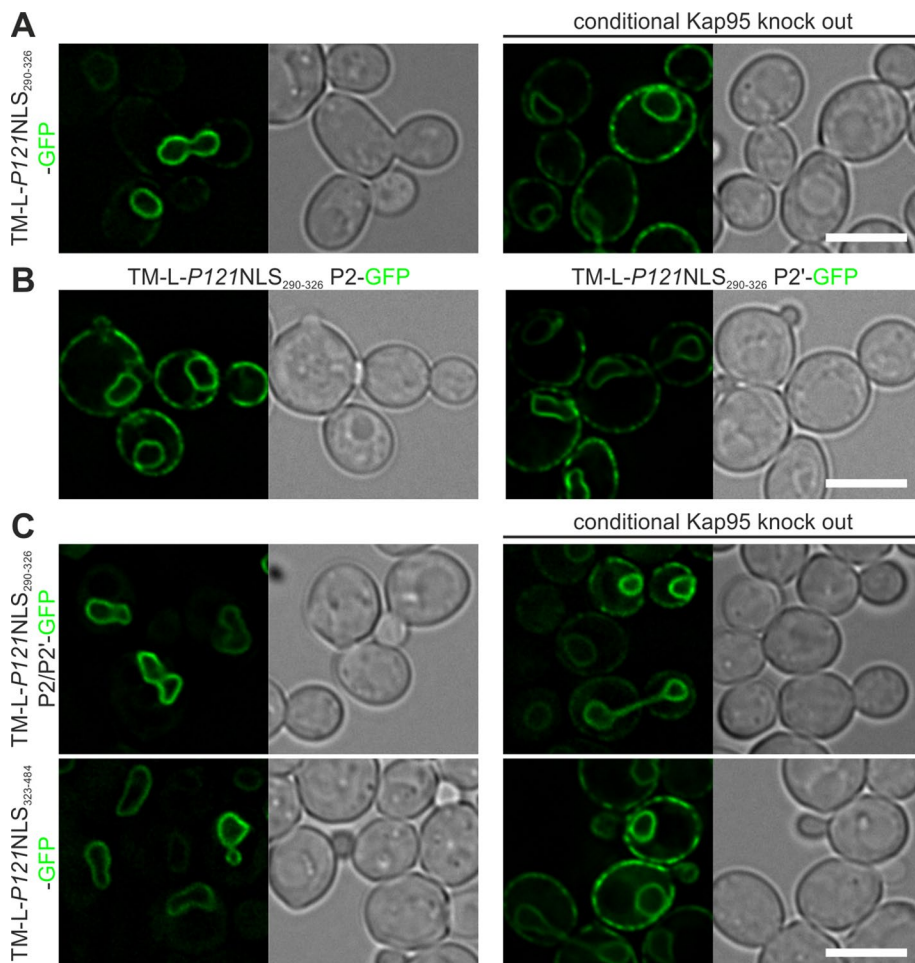


FIGURE 3: The IBB-like region is sufficient for INM targeting, and the *Pom121*NLS region consists of redundant NLSs. (A) The IBB-like region of *Pom121*NLS alone in TM-L-P121NLS₂₉₀₋₃₂₆-GFP is sufficient to induce Kap95-dependent INM accumulation of the reporter protein, and (B) substitution of the lysine residues at the P2 or P2' positions by alanine residues is sufficient to abolish NE accumulation. (C) Lysine-to-alanine mutation at the P2 and P2' positions of the IBB-like region of the NLS of *Pom121* in the context of the full *Pom121*NLS cluster (TM-L-P121NLS₂₉₀₋₄₈₄ P2/P2'-GFP) or removal of the IBB-like region from the full *Pom121*NLS (TM-L-P121NLS₃₂₃₋₄₈₄-GFP) did not alter Kap95-dependent INM accumulation of the indicated reporter proteins. Scale bars, 5 μ m.

***h2*NLS, *Pom121*NLS₂₉₀₋₄₈₄, and *Pom121*NLS₂₉₀₋₃₂₆ mediate nuclear import of a soluble reporter in HEK293T cells**

Thus far, our *in vivo* experiments in yeast reveal that the NLS of *Pom121* and the NLS of *Heh2* are largely interchangeable. To investigate whether this is also true in mammalian cells, we first performed a localization study in HEK293T cells with a soluble tandem GFP fused to the *Pom121*NLSs and *h2*NLS and mutant versions thereof (Figure 5). A tandem GFP (2GFP) protein localized dispersed over the cytoplasm and the nucleoplasm, and fusion to *h2*NLS, *Pom121*NLS₂₉₀₋₄₈₄, or *Pom121*NLS₂₉₀₋₃₂₆ caused strong accumulation in the nucleoplasm. The tandem GFP fused to the nucleophosmin NLS (2GFP-NPNLS) was used as a control. Mutant analysis showed that the accumulation of 2GFP with the *h2*NLS depends more strongly on the lysine at position P2' than on the residues at P2: ~30–40% of the cells expressing the P2' mutant showed cytoplasmic fluorescence, whereas for the wild type and the P2 mutant those percentages were >95%. This dependence on the residues at P2' rather than P2 was also found in yeast (Lokareddy *et al.*, 2015). A similar mutant analysis of the

*Pom121*NLS₂₉₀₋₄₈₄ showed that introduction of alanine residues at the P2 and P2' positions did not affect the localization in *Pom121*NLS₂₉₀₋₄₈₄. On the substitution of lysine by alanine at position P2 of 2GFP-*Pom121*NLS₂₉₀₋₃₂₆, the accumulation significantly decreased, resulting in 2GFP-NLS residing in the cytoplasm in all cells, whereas replacement of the P2' did not alter the localization of 2GFP-*Pom121*NLS₂₉₀₋₃₂₆. However, the P2' substitution was additive to the effect of P2 substitution in *Pom121*NLS₂₉₀₋₃₂₆ (2GFP-*Pom121*NLS₂₉₀₋₃₂₆ P2/P2') and resulted in complete lack of accumulation of 2GFP in the nucleoplasm in all cells. The fact that P2/P2' substitutions in *Pom121*NLS₂₉₀₋₄₈₄ did not influence this accumulation, whereas it does so in *Pom121*NLS₂₉₀₋₃₂₆, is consistent with previous studies (Doucet *et al.*, 2010; Yavuz *et al.*, 2010; Funakoshi *et al.*, 2011) and our conclusions from the localization studies in yeast that the *Pom121*NLS region has redundant importin α binding sites.

We conclude that *h2*NLS, *Pom121*NLS₂₉₀₋₄₈₄, and *Pom121*NLS₂₉₀₋₃₂₆ are all able to mediate nuclear accumulation of soluble tandem GFP in HEK293T cells, but there are different dependences of the interactions with the major and minor binding pockets of importin α . Whereas for *h2*NLS the interaction of the residue at P2' with importin α contributes mostly to efficient INM import, for *Pom121*NLS₂₉₀₋₃₂₆, the interaction of the P2 position with importin α is dominant for effective translocation of a soluble protein to the nucleus.

***h2*NLS targets a membrane protein to the INM in HEK293T cells**

We established that the *h2*NLS and *Pom121*NLS₂₉₀₋₃₂₀ have a similar IBB-like

interaction with importin α —that is, both target a soluble protein to the nucleus in HEK293T cells—and that the *Pom121*NLS mediates nuclear import of reporter and full-length membrane proteins in yeast. We next asked whether *h2*NLS is able to mediate INM import of a membrane-embedded reporter protein to the nucleus in mammalian cells. To answer this, we expressed GFP-*h2*NLS-L-TM, encoding *Heh*₉₃₋₃₇₈, in HEK293T cells. The reporter protein GFP-*h2*NLS-L-TM was clearly enriched at the nuclear envelope in HEK293T cells (Figure 6A). This enrichment was absent in cells expressing the reporter lacking the *h2*NLS (GFP-L-TM), which was dispersed over the NE/ER network, demonstrating the importance of the NLS for NE enrichment. The length of the ID linker region also influenced the localization of the reporter protein. A reporter with a shorter linker length of 37 instead of 180 residues, GFP-*h2*NLS-L(37)-TM, was enriched to the NE to some extent in a few cells, but in the majority of cells, this enrichment was minor or even absent, and the protein localized to the NE/ER network.

We wanted to know whether the NE enrichment of the reporter protein GFP-*h2*NLS-L-TM reflected INM localization of this

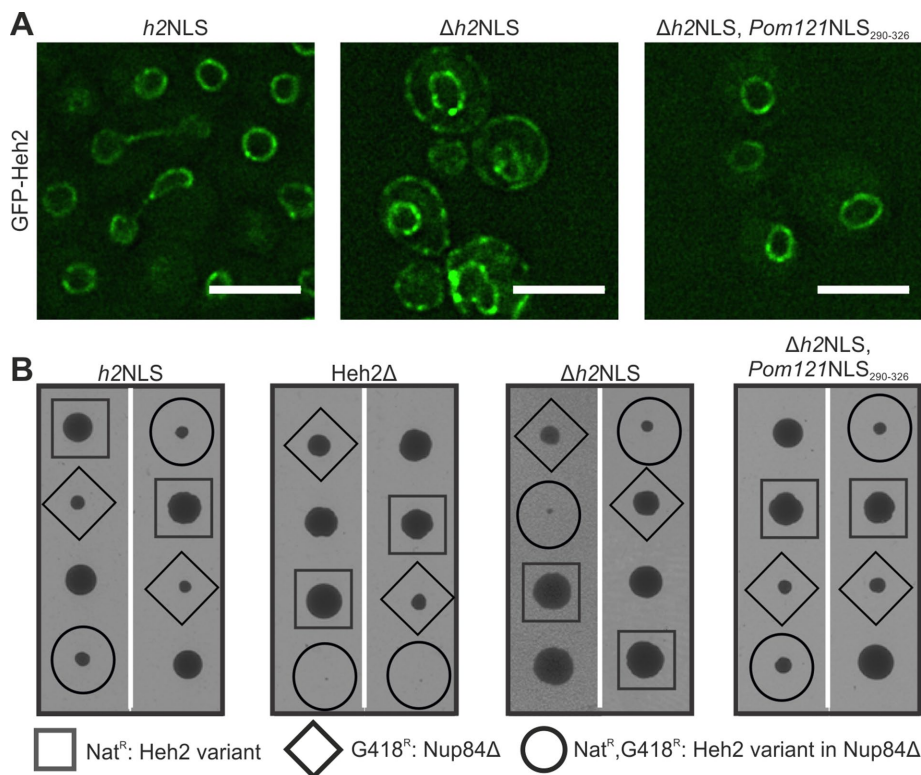


FIGURE 4: *Pom121NLS* rescues localization of *Heh2ΔNLS* and synthetic sickness of *Heh2ΔNLS,nup84Δ* double mutant. (A) Deconvolved wide-field images of yeast expressing native levels of GFP-Heh2 with wild-type NLS (*h2NLS*), without the NLS (*Δh2NLS*), or with *Pom121NLS*₂₉₀₋₃₂₆ instead of the *h2NLS* (*Δh2NLS, Pom121NLS*₂₉₀₋₃₂₆). Scale bar, 5 μm. (B) Synthetic sick/lethal interaction using tetrad dissection of *nup84Δ* expressing wild-type (*h2NLS*) and mutant variants (*Δh2NLS* and *Δh2NLS, Pom121NLS*₂₉₀₋₃₂₆) or no Heh2 (*Heh2Δ*). Each tetrad is oriented vertically and represents the meiotic progeny of a heterozygous diploid between GFP-HEH2-NAT/NUP84 and HEH2/*nup84::KANMX*. Two representative tetrads for each double mutant are shown. The genetic background of each spore is identified by the presence of the NAT or KAN marker, respectively. The double-mutant spore colonies are enclosed in circles, single mutants are enclosed in squares or diamonds, and wild-type strains are not enclosed.

membrane protein, and therefore we developed an assay based on the visualization of DNA methylation by an *Escherichia coli* DNA adenine methyltransferase (DAM) fused to the reporter protein: DAM-*h2NLS*-L-TM. DAM methylates adenine residues when it comes in contact with DNA. A GFP-fused truncation of *Dpn1* (GFP-Dpn7) specifically interacts with adenine-6-methylation (^m6A) in the sequence GATC (G^m6ATC; Kind et al., 2013). If an INM-localized DAM fusion methylates adenine residues at the nuclear periphery, like DAM fusion of lamin B1 does (Kind et al., 2013), this would be reflected in the specific localization of GFP-Dpn7 at the nuclear periphery. DAM-L-TM, lacking the *h2NLS*, served as the control. With a luminal domain of 54 kDa, this protein may be small enough to pass the NPC passively (Soullam and Worman, 1995; Wu et al., 2002; Ohba et al., 2004) but is not expected to accumulate at the INM because of lack of retention.

Indeed, when we coexpressed DAM-*h2NLS*-L-TM and GFP-Dpn7 in HEK293T cells, we observed increased fluorescence signal at the nuclear periphery of some cells (Figure 5B). Consistent with Kind et al. (2013), GFP-Dpn7 is homogeneously distributed over the nucleoplasm upon expression of DAM alone (unpublished data). However, localization of GFP-Dpn7 at the periphery was also observed in some cells expressing DAM-L-TM (Figure 6B), although the fraction of cells showing this GFP-Dpn7 localization is lower than for DAM-*h2NLS*-L-TM-expressing cells.

To determine whether there is a difference between cells expressing DAM-*h2NLS*-L-TM and DAM-L-TM, we determined the ratio of the average fluorescence signal at the nuclear periphery over the average fluorescence signal in the nucleoplasm (nuclear periphery vs. nucleoplasm, or P/N ratio). We plotted this ratio against the mean nuclear fluorescence signal of GFP-Dpn7 (Supplemental Figure S2A) and observed a correlation between the expression level of GFP-Dpn7 and the P/N ratio. Whereas a clear accumulation of GFP-Dpn7 at the nuclear periphery could be measured in cells with a low GFP-Dpn7 expression level, this accumulation was practically absent in cells with a high expression level. GFP-Dpn7 has a low affinity for unmethylated DNA, and therefore we reasoned that upon high expression of GFP-Dpn7, the DNA reaches a point of GFP-Dpn7 binding saturation, losing the resolution to determine the effect of the reporter-induced DNA methylation at the periphery. Therefore for our analysis we included only cells that have an average nucleoplasmic GFP-Dpn7 signal <90 (arbitrary units), using the same microscope settings between experiments. For cells transfected with DAM-L-TM, no significant difference was observed between P/N ratios of low and high GFP-Dpn7-expressing levels, whereas this difference was significant for cells transfected with DAM-*h2NLS*-L-TM (Supplemental Figure S2B). For cells expressing low levels of GFP-Dpn7, the mean P/N ratio was significantly higher than in cells expressing DAM-*h2NLS*-L-TM compared with cells expressing DAM-L-TM (Figure 6B). Because the P/N ratio is a measure for accumulation of GFP-Dpn7 at the nuclear periphery, we conclude that the NE enrichment that we observed for GFP-*h2NLS*-L-TM indeed reflects NLS-dependent INM localization of this reporter membrane protein.

DISCUSSION

With the discovery of active import of the membrane proteins Heh1 and Heh2 (King et al., 2006; Meinema et al., 2011) to the INM in baker's yeast, a main question was whether active transport of membrane proteins is yeast specific or conserved in higher eukaryotes. The domain composition of an NLS followed by an ID linker and a transmembrane domain, NLS-L-TM, is required and sufficient to mediate importin α/β-, FG-Nup-, and RanGTP-dependent import of Heh2 and Heh1 in baker's yeast (Meinema et al., 2011). Indeed, except for Heh1 and Heh2, no other INM proteins have been reported to traffic to the INM by an active import mechanism.

Here we screened metazoan genomes for genes that encode a putative NLS-L-TM domain composition. We identified four candidates—Sun2, Lem2, LBR, and Lap2β—and one for which the reversed topology (TM-L-NLS) was predicted, Pom121. The (putative) NLS-L or NLS regions of Sun2, Lem2, LBR, and Lap2β were not sufficient to mediate INM targeting of membrane reporter

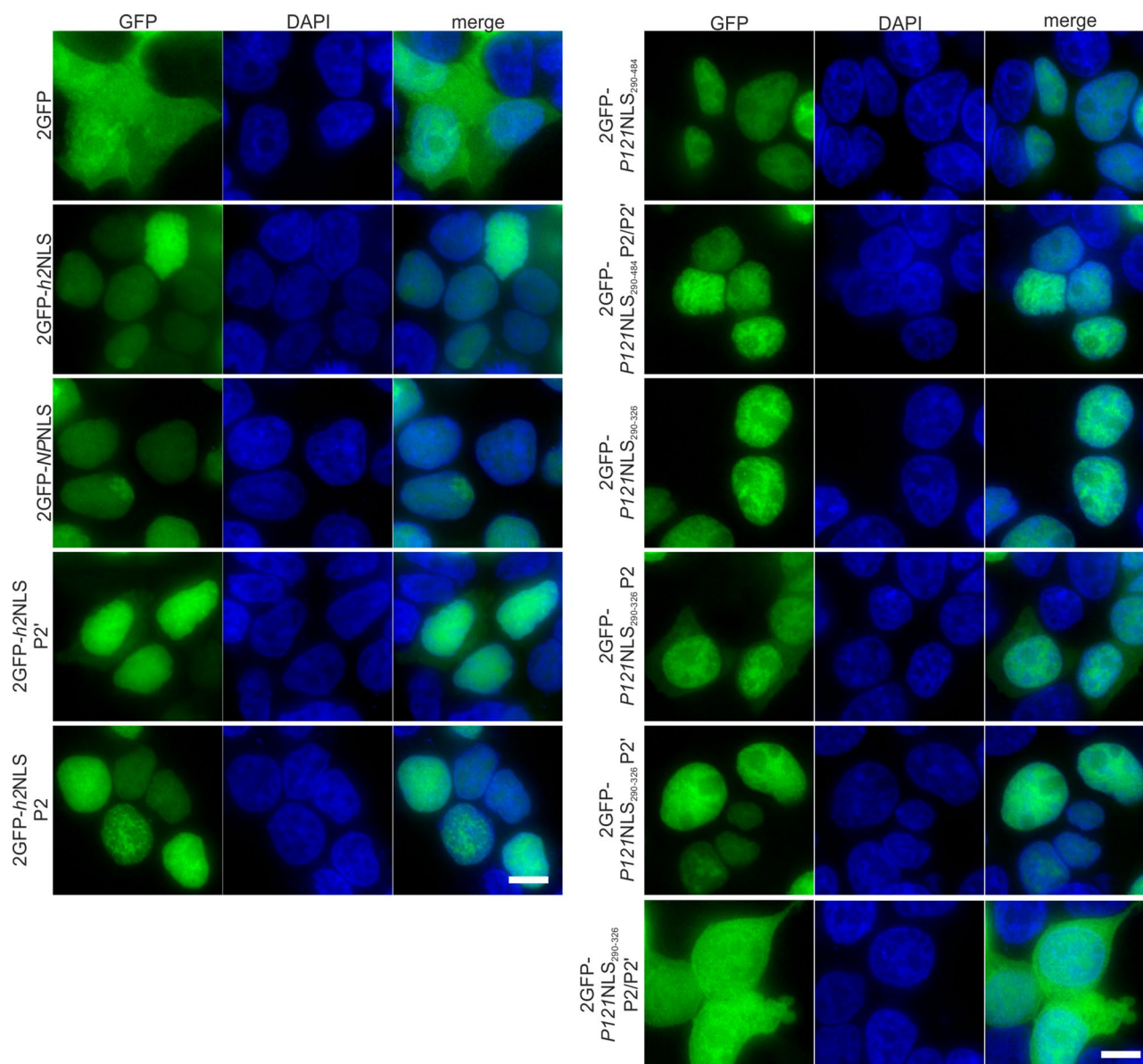


FIGURE 5: Localization of different GFP-NLS fusions in HEK293T cells. A tandem GFP fusion (2GFP; (top left) and 2GFP-Pom121NLS₂₉₀₋₃₂₆ P2/P2' (bottom right) equilibrate between the cytoplasm and the nucleoplasm. All remaining 2GFP-NLS fusion proteins accumulated in the nucleoplasm in >95% of the cells, with the exception of 2GFP-h2NLS P2'– and 2GFP-Pom121NLS₂₉₀₋₃₂₆ P2–expressing cells, in which cytoplasm-localized protein was observed in 30–40 and 100% of cells, respectively (100 < *n* < 200 cells). NPNLS: nucleoplasmin NLS. Scale bar, 5 μ m.

proteins in baker's yeast, but replacement of the h2NLS in TM-L-h2NLS-GFP for the NLS region of Pom121 (Pom121NLS₂₉₀₋₄₈₄) resulted in efficient accumulation at the INM, which depended on Kap60/95. The inability to mediate INM accumulation in yeast for the “NLS regions” of Sun2, Lem2, LBR, and Lap2 β obviously does not contradict their interaction with importin α and/or β , as previously demonstrated for LBR (Ma *et al.*, 2007) and Sun2 (Turgay *et al.*, 2010; Tapley *et al.*, 2011), and these signals could still promote INM localization but require mammalian-specific factors (such as lamins) to be retained once they reach the INM. Their insufficiency to drive accumulation in yeast merely categorizes them as being distinct from the Heh1, Heh2, and Pom121 NLSs, which are sufficient to support active import and accumulation of a membrane protein in yeast.

These experiments identify the Pom121 NLS as an NLS that supports active import of a membrane protein in yeast and also for the first time show that the NLS and linker regions can be N- or C-terminal of the transmembrane segment. We show that, as is the case for a C-terminal sorting signal, import depends on an NLS that interacts with importin α/β separated from a transmembrane domain by an intrinsically disordered region of sufficient length. This argues against the option that active import is related to the biogenesis of tail-anchored proteins and that membrane insertion may occur after nuclear import.

Having identified the NLS of Pom121 as an NLS that supports active import of a membrane protein in yeast, we address its similarities with the Heh1 and Heh2 NLSs. Crystallographic analysis of the binding interface of the Pom121NLS with importin α revealed

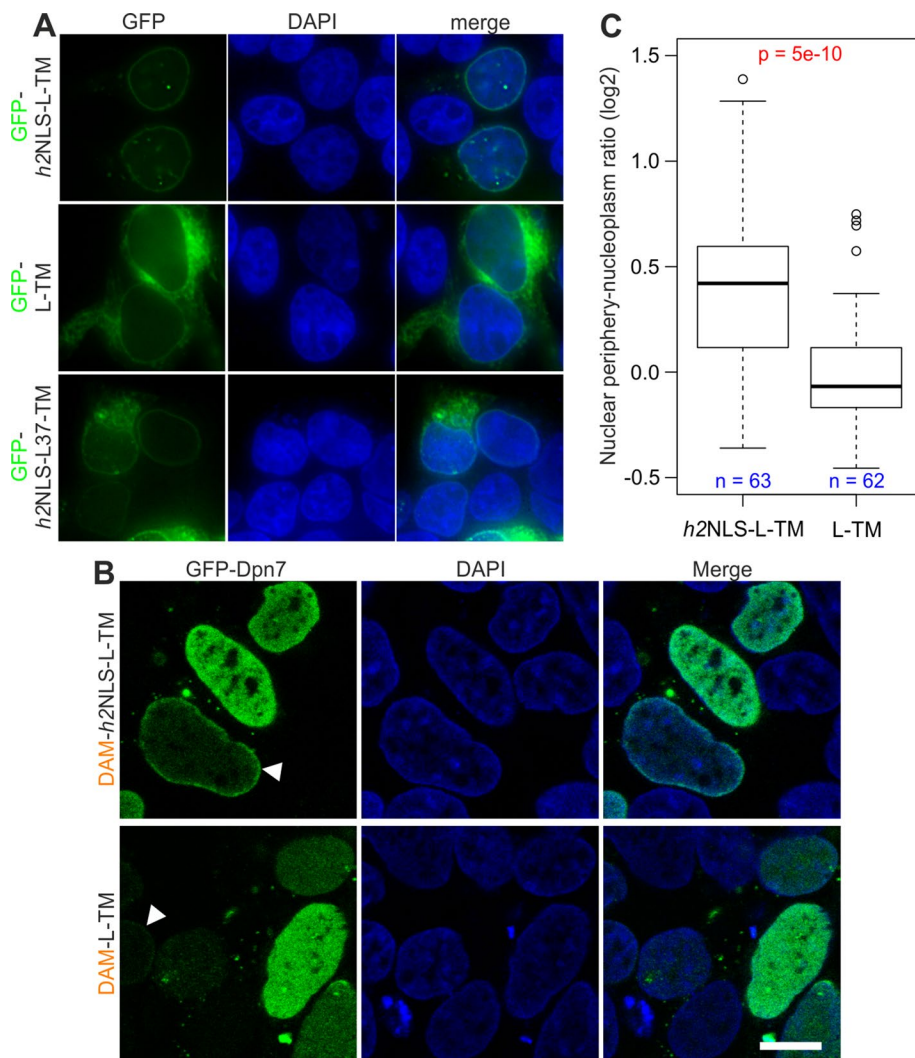


FIGURE 6: h2NLS mediates INM localization of a membrane-embedded protein in HEK293T. (A) The NE accumulation observed for GFP-h2NLS-L-TM is absent for GFP-L-TM, whereas GFP-h2NLS-L(37)-TM accumulates at the NE in a fraction of cells, and in the majority of cells, the reporter is NE/ER localized. Scale bar, 5 μ m. (B) Localization of GFP-Dpn7 in HEK293T expressing DAM-h2NLS-L-TM or DAM-L-TM. Arrowheads point at cells where GFP-Dpn7 accumulates at the nuclear periphery. Scale bar, 5 μ m. (C) Analysis of indicated number of cells with low GFP-Dpn7 levels shows that the accumulation of GFP-Dpn7 at the nuclear periphery, represented by the ratio of mean fluorescence signal at the nuclear periphery over the mean fluorescence signal in the nucleoplasm, is significantly lower for cells expressing DAM-L-TM than for cells expressing DAM-h2NLS-L-TM. The p values are calculated using Student's t test.

that of the five boxes of positively charged residues (Funakoshi *et al.*, 2011) in the NLS region (residues 290–484), the first two (P1'–P4' [294-KKKR-297] and P2–P5 [313-KRRR-316]) adopt a similar fold as the IBB domain of importin β bound to importin α . The NLS encoded by Heh1 and Heh2 adopts an IBB-like fold with intimate interactions in the minor and major binding sites (Lokareddy *et al.*, 2015). The tight interactions of Pom121NLS₂₉₁₋₃₂₀ with importin α are important for INM targeting, as mutagenesis of the residues interacting with the P2 and P2' positions of the Kap60 binding site in the Pom121NLS₂₉₀₋₃₂₆ (K313 for binding P2 or K295 for binding P2') results in lack of NE accumulation in yeast. These mutant NLSs are strong enough to accumulate a soluble cargo in human cells, hinting that the IBB-like features of the NLS may present a unique requirement for supporting INM import as compared with soluble import. The importance of the h2NLS for Heh2's function at the INM is clear

from the mislocalization and synthetic sickness of the double mutant Nup84 Δ , Heh2 Δ h2NLS. Additional proof for the conserved properties of the h2NLS and the Pom121NLS₂₉₀₋₃₂₆ come from experiments in which we assess their functionality in the context of native levels of Heh2. The subcellular localization of Heh2 in yeast is unaltered when its NLS is replaced for the Pom121NLS₂₉₀₋₃₂₆, and in addition, the Pom121NLS suppresses the synthetic sickness of Nup84 Δ , Heh2 Δ h2NLS.

Pom121 is the first metazoan INM protein identified to have a Heh1/Heh2/IBB-like NLS. The shared properties could be related to several factors, one being a shared role in nuclear import, which we investigated. We expressed yeast-derived reporter proteins in HEK293T cells to minimize effects of selective retention on localization, so that the readout of our assays (steady-state subcellular localization) would most likely reflect the dynamic equilibrium between rates of import and efflux. We observed clear NLS-dependent enrichment of the proteins at the NE. The dependence of linker length is less pronounced than in yeast, for which no accumulation is observed with a short linker of 37 residues. To confirm that the increased concentration at the NE indeed reflected an accumulation at the INM, we adapted an assay based on the visualization of methylated DNA at the nuclear periphery and showed that the NE accumulation indeed reflected the presence of DAM fusions of the membrane proteins at the INM.

Taken together, these data confirm NLS-dependent residence of the membrane proteins at the INM in HEK293T cells. However, when human cells are used, it is uncertain whether proteins traffic through the NPC or become INM localized during mitosis when the NE is reassembled onto the decondensing chromatin. Given the strong structural and biochemical similarities between the NLS of Heh1, Heh2, and Pom121, the ability

of Pom121NLS to support INM import in yeast, and the NLS-dependent localization of membrane proteins in HEK293T cells, we conclude that the mechanism is likely conserved for Pom121 but not for LBR, Sun2, and Lap2 β .

Two very recent studies (Boni *et al.*, 2015; Ungricht *et al.*, 2015) report on the determinants for membrane targeting in mammalian cells using LBR, Sun2, and Lap2 β as model substrates. The major determinants are the number and permeability of the NPCs, availability of binding sites at the INM, and kinetics of diffusion through the membranes of ER. Both studies convincingly show that a diffusion-retention model of INM protein transport in mammalian cells explains the measured kinetics of targeting in wild-type and mutant cells and under conditions of energy depletion. We here show that the NLSs in Heh1, Heh2, and Pom121 are distinct from the sorting signals in LBR, Sun2, and Lap2 β and therefore suggest that

targeting of Heh1, Heh2, and Pom121 does not follow this paradigm but instead is importin dependent, with higher tolerance for large extralumenal domains.

Although our genetic studies confirmed for the first time the *in vivo* importance of INM sorting of Heh2, the exact function of active import of Heh1, Heh2, and Pom121 remains to be determined. Intrigued by the elite position of Pom121, Heh1, and Heh2 in this respect, we speculate about why specifically these three proteins may require an active transport mechanism. A first consideration is that these proteins have large extralumenal domains, which may preclude fast enough passage through the lateral channels. Passive diffusion of Pom121 through the NPC is also less likely because of the FG repeats in its C-terminus, which promote interactions with components of the NPC. A second consideration is that the property of Pom121, Heh1, and Heh2 to accumulate efficiently at the INM might be essential for their role in the synthesis of new, functional pores. Heh1 and Heh2 are not stable components of the NPC, but they are involved in the biogenesis of intact NPCs in the NE (Yewdell *et al.*, 2011; Webster *et al.*, 2014). The synthetic sickness of the double mutant lacking Nup84 and expressing Heh2 Δ NLS may be related to the role of Heh2 in surveillance of NPC assembly (Webster *et al.*, 2014). Pom121 is dispensable for the assembly of NPCs in emerging NEs after mitosis (Funakoshi *et al.*, 2011), when “prepores” are seeded on mitotic chromosomes (Rasala *et al.*, 2008), but during interphase, Pom121 is essential for the insertion of new pores into the intact NE membrane (Doucet *et al.*, 2010; Talamas and Hetzer, 2011). To play an early role in NPC formation, Pom121 should not only localize to NPCs, but also reside at the INM. Indeed, a fraction of Pom121 is targeted to regions in the INM lacking intact NPCs (Funakoshi *et al.*, 2011). It was shown previously that Pom121 contains several bipartite NLSs, which interact with importin α/β and are required for efficient targeting of the protein to the NE/NPC (Doucet *et al.*, 2010; Yavuz *et al.*, 2010; Funakoshi *et al.*, 2011), although postmitotic NPC insertion was independent of the presence of the NLSs (Funakoshi *et al.*, 2011).

The identification of the interchangeability of the Pom121, Heh1, and Heh2 NLSs for nuclear traffic in yeast and humans cells provides the first experimental evidence in support of an active import mechanism to the INM in metazoans and warrants future investigation into its biological function.

MATERIALS AND METHODS

Molecular cloning

All plasmids are constructed according to standard molecular cloning techniques and listed in Supplemental Tables S3 and S5. All constructs generated in this study were fully sequenced to ensure the correctness of the DNA sequence. Details can be provided on request. In all constructs used in this study, the linker sequences originated from Heh2, the sequence encoding the TM in constructs with topology N-TM-C_{lumen} originated from TM1 from Heh2, and the sequence encoding the TM in constructs with topology N_{lumen}-TM-C was encoded by a synthetic gene (encoding a reversed orientation of the amino acids of Heh2 TM). The exact encoded amino acid sequences and their origin are provided in Supplemental Tables S6 and S7. The gene encoding human importin α 1 was cloned as FL and Δ IBB in vectors pET28a (Novagen, Merck Millipore, Darmstadt, Germany) and pGEX-6P (GE Healthcare) as previously described (Pumroy *et al.*, 2015). Murine Δ IBB-importin α 1 used for crystallization was expressed and purified as described (Lott *et al.*, 2010). The NLS sequence of Rn Pom121 (residues 291–320) was cloned between *Bam*HI and *Xho*I restriction sites in vectors pGEX-6P and in an engineered pET28 that also contains an N-terminal MBP (pET28-MBP). NPNLS and h2NLS were also cloned in pET28-MBP as previously

described (Lokareddy *et al.*, 2015). Alanine mutations at position P2, P2', and P2/P2' of pET28-MBP- Pom121NLS₂₉₁₋₃₂₀ were generated by site-directed mutagenesis.

Yeast cultivation

All reporter proteins were expressed in *S. cerevisiae* KAP95AA strain (Meinema *et al.*, 2011). Strains were grown at 30°C in synthetic dropout medium lacking histidine and supplemented with 0.01% adenine and 2% glucose. For induction of the GAL1 promoter, glucose was replaced by raffinose and 0.1% galactose was added for 2 h; full-length expression was confirmed by Western blot (Supplemental Figure S3).

Depletion of cytosolic Kap95-FRB was induced by the addition of 2 μ g/ml rapamycin to the cell culture for 15 min.

Synthetic lethality screening

DNA constructs encoding GFP-tagged, full-length Heh2, GFP-tagged Heh2(Δ h2NLS), and GFP-tagged Heh2(Δ h2NLS, Pom121NLS₂₉₁₋₃₂₀) were fused to the NAT marker and the flanking regions as encoded immediately upstream and downstream the HEH2 open reading frame. The linear DNA constructs were subsequently integrated in BY4742 *heh2::KAN* by homologous recombination, thereby replacing the KAN marker for GFP-HEH2-NAT (Supplemental Table S4). Transformants were selected and analyzed for expression of the GFP-tagged proteins by fluorescence microscopy. For assessing the synthetic sick and lethal phenotypes of double mutants, BY4741 *nup84::KAN* (Mata) was mated with the GFP-Heh2 variant expressing strains AS1 (GFP-HEH2-NAT, Mat α), AS2 (GFP-HEH2(Δ h2NLS)-NAT, Mat α), AS5 (GFP-HEH2(Δ h2NLS, Pom121NLS₂₉₁₋₃₂₀)-NAT, Mat α), and AS4 (GFP-NAT, Mat α). Diploids were selected on yeast extract/peptone/dextrose (YPD) containing G418 and Nat. After sporulation, tetrads were dissected, and the genotype of each spore was determined by replica plating on YPD containing Nat and YPD containing G418.

Transfection HEK293T cells for microscopy

HEK293T cells were grown in DMEM (Invitrogen) supplemented with antibiotics and 10% fetal calf serum at 37°C and 5% CO₂. Cells were grown on coverslips and transiently transfected using FuGENE HD Transfection reagent (Promega). Subsequently the cells were fixed with 4% paraformaldehyde in phosphate-buffered saline (PBS), permeabilized with 0.1% Triton X-100 in PBS, stained with 1 μ g/ml 4',6-diamidino-2-phenylindole (DAPI), and mounted with Vectashield (Vector Labs, Burlingame, CA).

Fluorescence microscopy

Imaging of localization of GFP-fused reporter proteins in yeast and HEK293T cells was done on a DeltaVision Deconvolution Microscope (Applied Precision), using InsightSSI Solid State Illumination of 488 nm (GFP) and 358 nm (DAPI). Detection was done with a CoolSNAP HQ² camera. For yeast, an Olympus UPLS Apo 100 \times oil objective with 1.4 numerical aperture (NA) was used. Pixel size was 64 \times 64 \times 200 nm. For HEK293T an Olympus UPLS Apo 40 \times oil objective with 1.3 NA was used. Pixel size was 215 \times 215 \times 500 nm.

DAM methylation assay

HEK293T cells were cultured as described. Cells were cotransfected with 0.8 μ g of plasmid encoding DAM-fused reporter proteins (pIND-DAM-V5 (Vogel *et al.*, 2006), pAK49, and pAK50, respectively), 0.8 μ g of enhanced GFP-DPN7 (Kind *et al.*, 2013), and 0.8 μ g of pVg-RxR (Invitrogen) using 7.5 μ l of TransIT-2020 transfection reagent (Mirus) for 24 h. Expression of DAM-fusion proteins was induced by the addition of 5 μ M ponasterone (Sigma-Aldrich, St. Louis, MO) for

24 h. Cells were fixed with 4% paraformaldehyde, permeabilized with 0.1% Triton X-100 and 0.5% bovine serum albumin, stained with 1 µg/ml DAPI, and mounted with Vectashield.

Imaging of eGFP-Dpn7 and DAPI was done on a Leica TCS SP5 laser scanning confocal microscope using a 488-nm argon gas laser for GFP, a 405-nm diode laser for DAPI, and a 63×/1.4 NA oil immersion objective. Detection was performed with a photomultiplier tube. Bidirectional line scanning was used with a line frequency of 700 Hz, and four frames were averaged. Settings were kept constant. Data were analyzed using the ZEN2010B software package (Carl Zeiss, Jena, Germany).

Biochemical techniques

FL- and Δ IBB-importin α 1 was expressed and purified as previously described (Lott *et al.*, 2011; Pumroy *et al.*, 2015). The Δ IBB-importin α 1:*Pom121NLS*₂₉₁₋₃₂₀ complex was formed by coexpressing plasmids pET28a- Δ IBB-importin α 1 and pGEX-*Pom121NLS*₂₉₁₋₃₂₀ in *E. coli* strain BL21-CodonPlus (DE3)-RIL (Stratagene) for 6 h at 30°C. GST-*Pom121NLS*₂₉₁₋₃₂₀ bound to Δ IBB-importin α 1 was purified on glutathione beads (GenScript), and after cleaving off the GST with PreScission Protease, the complex was purified over a Superdex 200 column (GE Healthcare) equilibrated in G.F. buffer (20 mM Tris, pH 8.0, 150 mM NaCl, 5 mM β -mercaptoethanol, and 0.2 mM phenylmethylsulfonyl fluoride). All GST-tagged constructs used in this study were purified as described earlier. All MBP-tagged NLSs (*Pom121NLS*₂₉₁₋₃₂₀, NP-NLS, h2NLS) were expressed and purified as by Pumroy *et al.* (2015) and Lokareddy *et al.* (2015). The IBB-displacement assay on glutathione beads in Figure 2 was carried out and quantified as previously described (Lokareddy *et al.*, 2015; Pumroy *et al.*, 2015). The error bars in the quantification represent the SD of three independent experiments carried out under identical conditions.

Crystallographic studies

Crystals of Δ IBB-importin α 1 bound to *Pom121NLS*₂₉₁₋₃₂₀ were obtained by mixing equal volume of gel filtration-purified complex at 15 mg/ml with 0.1 M sodium citrate tribasic dihydrate, pH 5.6, 0.7 M sodium citrate tribasic dihydrate, and 25 mM dithiothreitol and equilibrating the droplet against 600 µl of the same precipitant. Crystals were harvested in nylon cryoloops, cryoprotected with 27% ethylene glycol, and flash-frozen in liquid nitrogen. Crystals were diffracted at beamlines X6A and X29 at the National Synchrotron Light Source (Brookhaven National Laboratory, Upton, NY) on a Quantum Q270 and a Quantum-315r charge-coupled device detector, respectively. Data indexing, integration, and scaling were carried out with the HKL2000 (Otwinowski and Minor, 1997). Initial phases were obtained by molecular replacement using Phaser (McCoy *et al.*, 2007) and Protein Data Bank entry 3Q5U as a search model. Atomic models were built using Coot (Emsley and Cowtan, 2004) and refined using *phenix.refine* (Adams *et al.*, 2002). A continuous trace of *Pom121NLS*₂₉₁₋₃₂₀ main chain between residues 299 and 307 was obtained by blurring the *B* factor by applying a positive *B* factor correction of 20 Å² in Coot. The refined *B* factor of this region is obviously very high (120 Å²). Data collection and refinement statistics are summarized in Supplemental Table S1. The structure was analyzed using the PISA server (Xu *et al.*, 2008) and PyMOL (PyMOL Molecular Graphics System, version 1.5.0.4; Schrödinger LLC).

Accession code

The atomic coordinates and structure factors for Δ IBB-importin α 1 bound to *Pom121NLS* were deposited in the Protein Data Bank with accession code 4YI0.

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Supplemental Materials

Molecular Biology of the Cell

Kralt et al.

Supplemental Material to:

Conservation of inner nuclear membrane targeting sequences in mammalian Pom121 and yeast Heh2 membrane proteins

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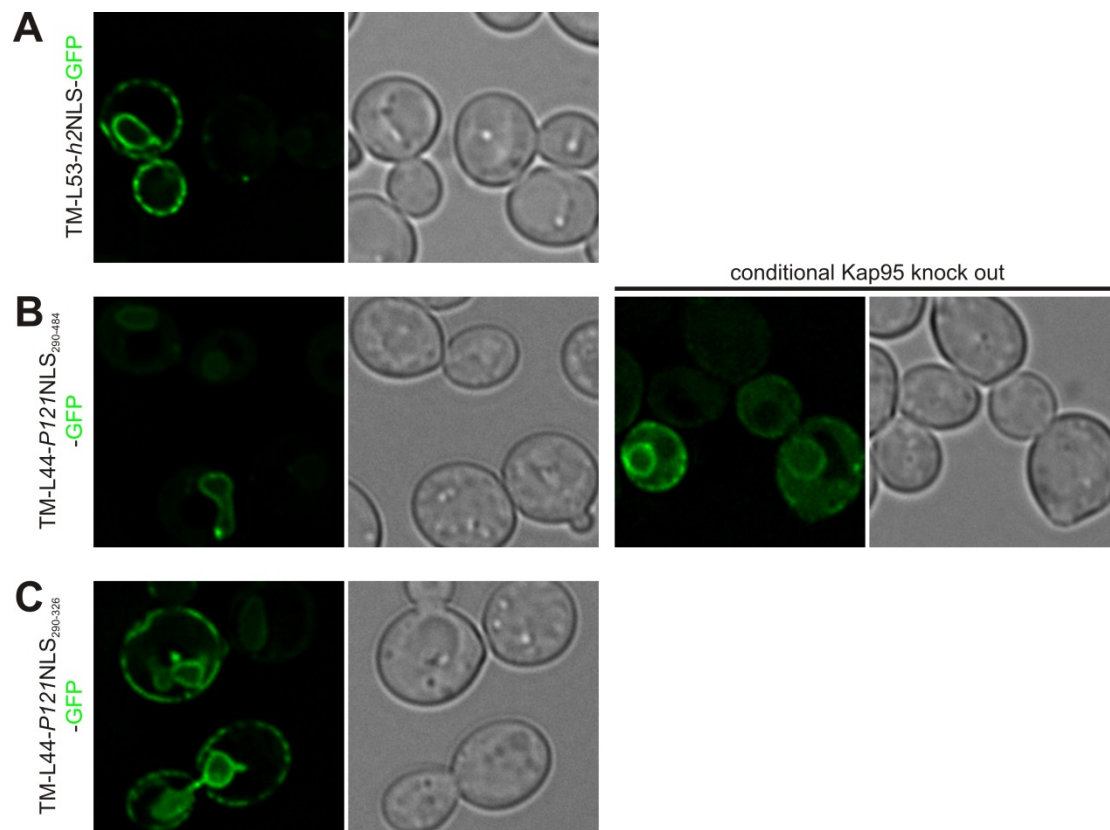


Figure S1 An ID linker region between the transmembrane segment and the NLS is required for INM import of proteins with TM-L-NLS topology (A) TM-L53-h2NLS-GFP, with an ID linker region that was shortened from 180 to 53 residues, did not accumulate at the INM, but was dispersed over the NE-ER network. (B) TM-L44-*Pom121NLS*₂₉₀₋₄₈₄-GFP with reduced ID linker region and the full *Pom121NLS* still localized to the INM in a Kap95-dependent manner. (C) TM-L44-*Pom121NLS*₂₉₀₋₃₂₆-GFP, which encodes the shortened ID linker and only the IBB-like region of *Pom121NLS*, localized to the NE-ER network and did not accumulate at the INM. The accumulation in (B) may be explained as the ~70 residues of the IBB-like and the spacing region to the third stretch of basic residues of the *Pom121NLS* region could function as part of the ID linker region, and the second C-terminal importin binding site in the *Pom121NLS* region mediates INM localization of this reporter.

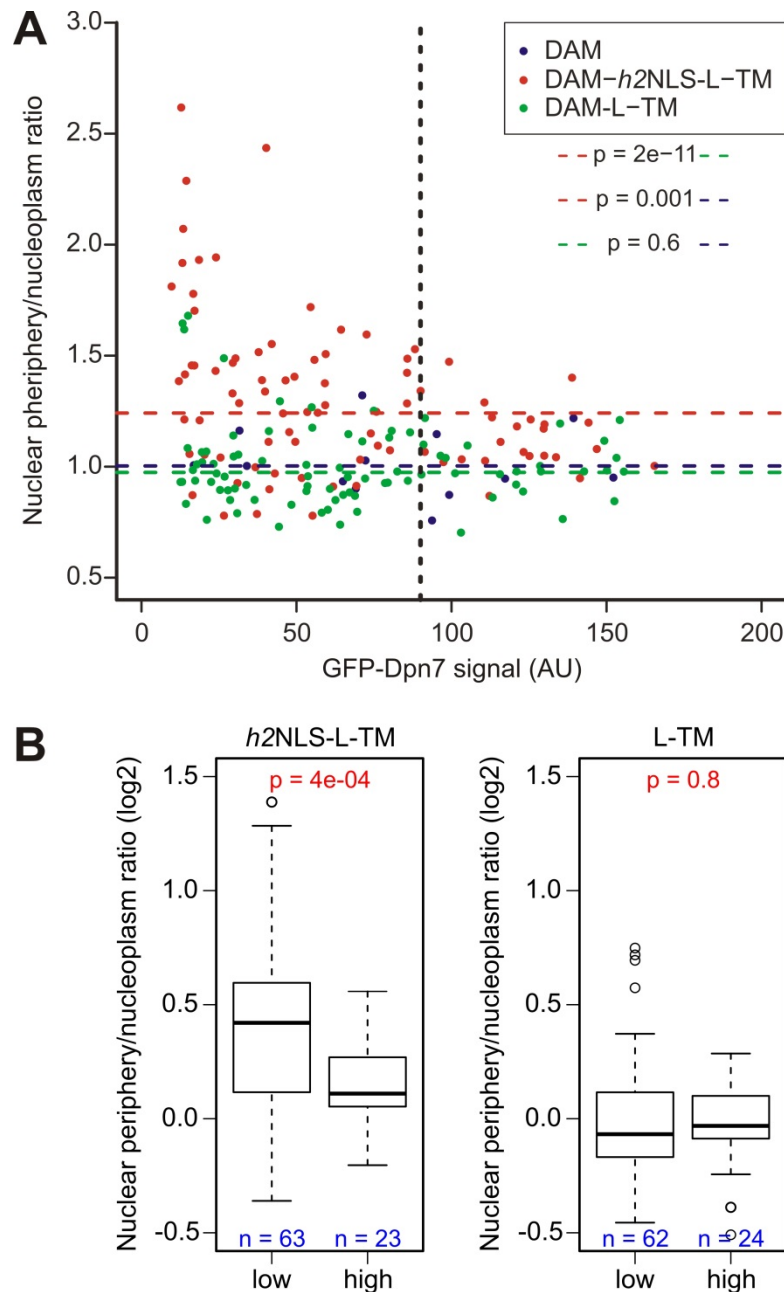


Figure S2 DAM methylation of DNA at the nuclear periphery confirms the NLS dependent INM localization of these proteins (A) The accumulation of GFP-Dpn7 at the periphery of the nucleus in cells expressing DAM (blue), DAM-*h2NLS-L-TM* (red) and DAM-*L-TM* (green) is represented as the ratio of average fluorescent signal at the cell periphery over the average fluorescent signal in the nucleoplasm (periphery-nucleoplasm ratio or P/N ratio). The periphery-nucleoplasm ratio was plotted against the average GFP-Dpn7 signal in the nucleus. The mean ratios are indicated by the dotted lines. An arbitrary cut-off between cells that express high and low levels of GFP-Dpn7 was set at 90 AU (black dotted line). P-values are calculated using the Wilcoxon test. (B) Boxplots comparing the P/N ratios of D-*h2NLS-L-TM* and D-*L-TM* expressing cells with low or high GFP-Dpn7 expression levels. No significant difference was found for cells expressing *L-TM*, but a significant difference was found for cells expressing *h2NLS-L-TM*. Logged data follows a normal distribution according to Shapiro-Wilk and P-values are calculated using Student t-test.

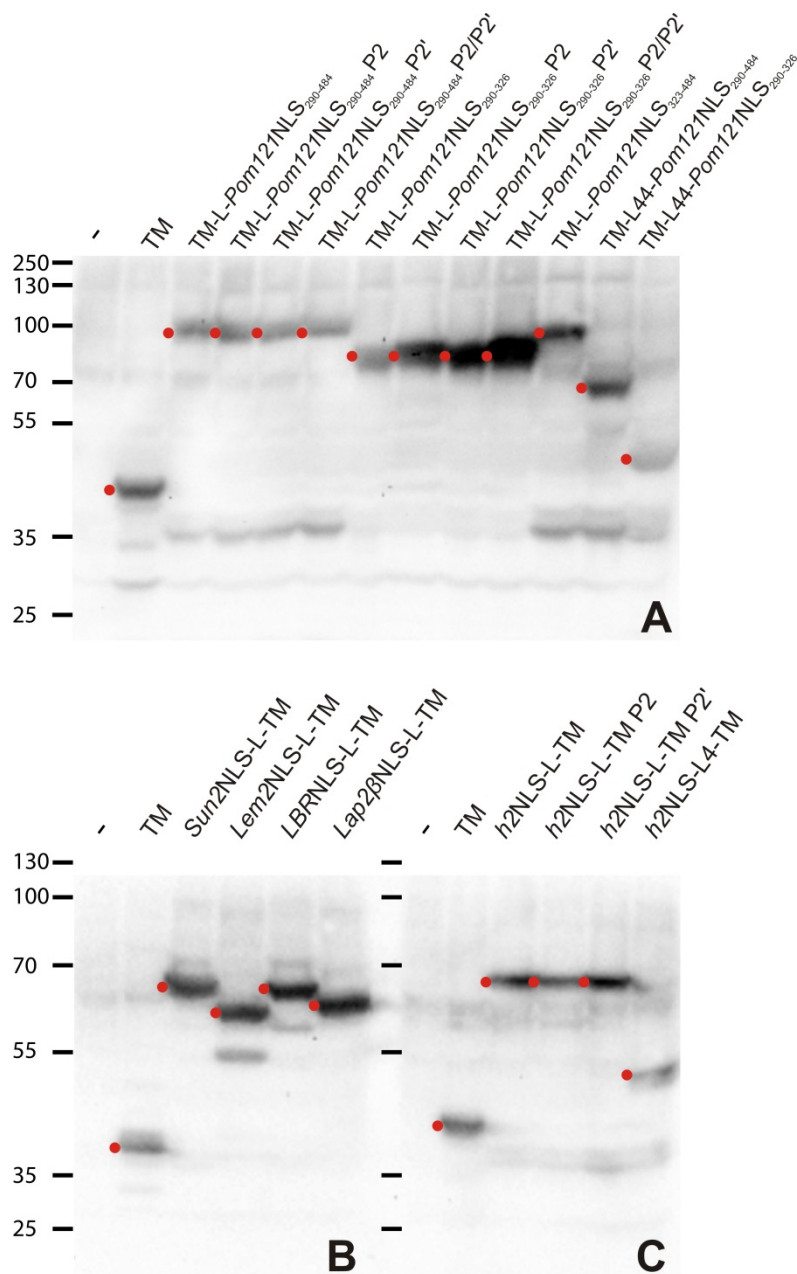


Figure S3 Visualization of expression of NLS-L-TM and TM-L-NLS reporter proteins in yeast by Western blotting. Cell lysates were prepared from equal amounts of *S.cerevisiae* KAP95AA cells expressing C-terminal (A,C) and N-terminal GFP-tagged reporter proteins and anti-GFP was used for immuno-detection.

Table S1: Data collection and refinement statistics.

ΔIBB-importin α1:Pom121NLS	
Data collection	
Space group	P2 ₁ 2 ₁ 2 ₁
Cell dimensions	
<i>a</i> , <i>b</i> , <i>c</i> (Å)	78.3, 89.8, 97.9
α , β , γ (°)	90.0, 90.0, 90.0
Resolution (Å)	15-1.80
No. reflections (tot/unique)	992,426 / 62,116
<i>R</i> _{sym}	5.7 (50.1)
<i>I</i> / σ <i>I</i>	34.7 (2.1)
Completeness (%)	97.8 (98.3)
Redundancy	3.6 (3.2)
Refinement	
Resolution (Å)	15-1.80
No. reflections used	60,036
<i>R</i> _{work} / <i>R</i> _{free}	18.2 / 20.9
No. atoms	
Protein	3,241
Ligand (Pom121NLS)	449
Water	607
<i>B</i> -factors (Å ²)	
Protein	28.6
Ligand/ion	71.8
Water	39.6
R.m.s. deviations	
Bond lengths (Å)	0.007
Bond angles (°)	1.09

Values in parentheses are for highest-resolution shell (1.86-1.80).

The *R*_{free} was calculated using 5% of randomly selected reflections.

Table S2: Structural alignment of Pom121NLS with other NLSs visualized crystallographically in complex with importin α and Kap60.

<u>NLS type</u>	<u>Minor Binding Site</u>	<u>Major Binding Site</u>	<u>PDB id</u>
	P ₋₁ 'P ₀ 'P ₁ 'P ₂ 'P ₃ 'P ₄ 'P ₅ '	P ₁ P ₂ P ₃ P ₄ P ₅	
SV40T-ag	P K K K R K	P K K K R K V	1EJL/1BK6*
hPLSCR1-NLS		G K I S K HWTGI	1Y2A
hPLSCR4-NLS	S I I R K W N		3Q5U
Gua α -NLS	G Q K R S F S		3ZIN
A89-NLS	L G K R K Y W		4B8P**
B54-NLS	L G K R K R H		2YNS**
TPX2	K R K H	P V K M I K	3KND
C-Myc	K R V K L	P A A K R V K	1EE4*
NP	A V K R P A A	TKKAGQ A K K K K L D	1EJY/1EE5*
Kap60-IBB	E L R R R R D	TQQVELRKAKRDEA L A K R R N F	1WA5*
h1NLS	TR K K R K D P	DSDDWSES N S K E N K ID	4XZR *
h2NLS	T N K R K R E	QISTDNEAKMQIQEEKS P K K K R K KRSSKANK	4PVZ*
Pom121NLS	K E K K K R T	VAEEDQLHLDGQ E N K R R R HDSS	4YI0*

* denotes yeast importin α (Kap60).

** denotes rice importin α . In all other cases, mammalian importin α was co-crystallized with NLSs α

Table S3: Plasmids for *S. cerevisiae*

Plasmid Name (Reporter name)	Encodes ³
pAK80 (GFP-TM) ¹	GFP-Heh2 ₃₀₀₋₃₇₉
pACM023 (GFP-h2NLS-L-TM) ¹	GFP-Heh2 ₉₃₋₃₇₉
pAK99 (<i>Sun2</i> NLS-L-TM) ¹	GFP- <i>HsSun2</i> ₁₈₋₁₀₆ -Heh2 ₁₃₈₋₃₇₉
pAK100 (<i>Lem2</i> NLS-L-TM) ¹	GFP- <i>MmLem2</i> ₁₂₈₋₁₄₅ -Heh2 ₁₃₈₋₃₇₉
pAK102 (<i>LBR</i> NLS-L-TM) ¹	GFP- <i>HsLBR</i> ₅₅₋₁₁₃ -Heh2 ₁₃₈₋₃₇₉
pAK103 (<i>Lap2β</i> NLS-L-TM) ¹	GFP- <i>HsLap2β</i> ₂₄₈₋₂₆₅ -Heh2 ₁₃₈₋₃₇₉
pAK01 (TM-GFP) ²	TM-GFP (encodes synthetic TM)
pAK54 (TM-L- <i>Pom121</i> NLS ₂₉₀₋₄₈₄) ²	TM-Heh2 ₁₄₅₋₃₁₅ - <i>Pom121</i> ₂₉₀₋₄₈₄ -GFP
pAK55 (TM-L- <i>h2</i> NLS) ²	TM-Heh2 _{145-315,93-145} -GFP
pAK95 (TM-L- <i>Pom121</i> NLS ₂₉₀₋₃₂₆) ²	TM-Heh2 ₁₄₅₋₃₁₅ - <i>Pom121</i> ₂₉₀₋₃₂₆ -GFP
pAK96 (TM-L- <i>Pom121</i> NLS ₂₉₀₋₃₂₆ _P2) ²	TM-Heh2 ₁₄₅₋₃₄₅ - <i>Pom121</i> ₂₉₀₋₃₂₆ (K313→A) -GFP
pAK97 (TM-L- <i>Pom121</i> NLS ₂₉₀₋₃₂₆ _P2') ²	TM-Heh2 ₁₄₅₋₃₄₅ - <i>Pom121</i> ₂₉₀₋₃₂₆ (K295→A) -GFP
pAK105 (TM-L- <i>Pom121</i> NLS ₂₉₀₋₄₈₄ _P2/P2') ²	TM-Heh2 ₁₄₅₋₃₄₅ - <i>Pom121</i> ₂₉₀₋₄₈₄ (K295 and 313→A) -GFP
pAK107 (TM-L- <i>Pom121</i> NLS ₃₂₃₋₄₈₄) ²	TM-Heh2 ₁₄₅₋₃₄₅ - <i>Pom121</i> ₃₂₃₋₄₈₄ -GFP
pAK93 (TM-L44- <i>Pom121</i> NLS ₂₉₀₋₄₈₄) ²	TM-Heh2 _{145-153,305-315} - <i>Pom121</i> ₂₉₀₋₄₈₄
pAK94 (TM-L53- <i>h2</i> NLS) ²	TM-Heh2 _{145-153,305-315,93-145}
pAK104 (TM-L44- <i>Pom121</i> NLS ₂₉₀₋₃₂₆) ²	TM-Heh2 _{145-153,305-315} - <i>Pom121</i> ₂₉₀₋₃₂₆

¹Originates from pUG34, *MET25* promoter is replaced by *GAL1* promoter, N-terminal GFP fusion, HIS3

²Originates from pUG23, *MET25* promoter is replaced by *GAL1* promoter, C-terminal GFP-fusion, HIS3

³In the Heh2 proteins residues 93-138 are encoding the NLS region and residues 138-379 encode the linker region plus the first TM region. Residues 145-315 encode a major part of the linker region.

Table S4: *S. cerevisiae* strains

Strains name	Genotype	Sources
HHY118 (<i>Kap95AA</i>)	MAT α tor1-1 frp1::NAT PMA1-2xFKBP12::TRP1 KAP95-FRB::KAN	(Haruki et al., 2008)
<i>heh2</i> Δ	BY4742 <i>heh2</i> ::KAN	Yeast KO collection
AS1	BY4742 <i>heh2</i> ::GFP-HEH2-NAT	This study
AS2	BY4742 <i>heh2</i> ::GFP-HEH2(Δ h2NLS)-NAT	This study
AS5	BY4742 <i>heh2</i> ::GFP-HEH2(Δ h2NLS, <i>Pom121</i> NLS ₂₉₁₋₃₂₀)-NAT	This study
AS4	BY4742 <i>heh2</i> ::GFP-NAT	This study
BY4741	MAT α <i>his3</i> Δ 1 <i>leu2</i> Δ 0 <i>met15</i> Δ 0 <i>ura3</i> Δ 0	Invitrogen
<i>nup84</i> Δ	BY4741 <i>nup84</i> Δ ::KAN	Yeast KO collection

Table S5: Plasmids for HEK293T cells

Plasmid Name (Reporter name)	Encodes ³
pAK111 (2GFP) ¹	2GFP
pAK112 (2GFP- <i>h2</i> NLS) ¹	2GFP-Heh2 ₉₃₋₁₄₅
pAK113 (2GFP- <i>h2</i> NLS_P2) ¹	2GFP-Heh2 ₉₃₋₁₄₅ (K126→A)
pAK114 (2GFP- <i>h2</i> NLS_P2') ¹	2GFP-Heh2 ₉₃₋₁₄₅ (R103→A)
pAK115 (2GFP- <i>Pom121</i> NLS ₂₉₀₋₄₈₄) ¹	2GFP-Pom121 ₂₉₀₋₄₈₄
pAK118 (2GFP- <i>Pom121</i> NLS ₂₉₀₋₄₈₄ P2/P2') ¹	2GFP-Pom121 ₂₉₀₋₄₈₄ (K295 and 313→A)
pAK119 (2GFP- <i>Pom121</i> NLS ₂₉₀₋₃₂₆) ¹	2GFP-Pom121 ₂₉₀₋₃₂₆
pAK120 (2GFP- <i>Pom121</i> NLS ₂₉₀₋₃₂₆ _P2) ¹	2GFP-Pom121 ₂₉₀₋₃₂₆ (K313→A)
pAK121 (2GFP- <i>Pom121</i> NLS ₂₉₀₋₃₂₆ _P2') ¹	2GFP-Pom121 ₂₉₀₋₃₂₆ (K295→A)
pAK122 (2GFP- <i>Pom121</i> NLS ₂₉₀₋₃₂₆ _P2P2') ¹	2GFP-Pom121 ₂₉₀₋₃₂₆ (K295 and 313→A)
pAK123 (2GFP-NPNLS) ¹	2GFP-X/Nucleoplasmin ₁₅₃₋₁₇₂
pAK37 (GFP- <i>h2</i> NLS-L-TM) ¹	GFP-Heh2 ₉₃₋₃₇₈
pAK38 (GFP-L-TM) ¹	GFP-Heh2 ₁₃₈₋₃₇₈
pAK108 (GFP- <i>h2</i> NLS-L37-TM) ¹	GFP-Heh2 _{93-145,289-378}
pAK49 (DAM- <i>h2</i> NLS-L-TM) ²	DAM-Heh2 ₉₃₋₃₇₈
pAK50 (DAM-L-TM) ²	DAM-Heh2 ₁₃₈₋₃₇₈

¹Originates from peGFP-c1

²Originates from pIND/V5-HisA (Invitrogen and (Vogel *et al.*, 2006)

Table S6: Amino acid sequences of reporter proteins in *S. cerevisiae*¹

GFP-TM (pAK80) (TM: Heh2 ₃₀₀₋₃₇₉)	MSKGEELFTGVVPILVELDGDVNGHKFSVSGEGEDATYGKLT IK FICTTGKLPVPWPTLVTTFGYGVQC FARYPDHMKQHDFFSKAMPEGYVQERTIFFKDDGNYKTRAEVKFEGDTLVNRIELKGIDFKEDGNILGHK LEYN Y NSHN V YIMADKQKNGIKVNF K IRHNIEDGSVQLADHYQQNTPIGDPVLLPDNHYLSTQSALS KD PNEKRDH M VLLFEVTAAGITHGMDELYKSRTSGSRMKANKTKRGIDIMKPFIAHLFIWLNWNGAIFLSIICP ILFGLWYREQRIQVGYCGHEKPLKSLAISAFPQTERVDSVLQAYRP
GFP-NLS-L-TM (L-TM: Heh2 ₁₃₈₋₃₇₉) (At “_NLS_” the below indicated NLS(-like) sequences are introduced)	MSKGEELFTGVVPILVELDGDVNGHKFSVSGEGEDATYGKLT IK FICTTGKLPVPWPTLVTTFGYGVQC FARYPDHMKQHDFFSKAMPEGYVQERTIFFKDDGNYKTRAEVKFEGDTLVNRIELKGIDFKEDGNILGHK LEYN Y NSHN V YIMADKQKNGIKVNF K IRHNIEDGSVQLADHYQQNTPIGDPVLLPDNHYLSTQSALS KD PNEKRDH M VLLFEVTAAGITHGMDELYKSRTS_NLS_PPESPPQSKSDGKATSADLTSELTVEELHKKDS SDDKPRVKELPKPELNLKVSNEFLAQLNKLASAATENYDHSIKSTDLSSIRIETEEPVGPSTGAETRNESE VMENINLEVQPEVKEAKEELTKISETFDNQDEEDTSRLSSKNIRSPKGRTRHFIANKTKRGIDIMKPFIAHL FIWLNWNGAIFLSIICPILFGLWYREQRIQVGYCGHEKPLKSLAISAFPQTERVDSVLQAYRP
<i>h2NLS</i> (pACM023) (<i>Saccharomyces cerevisiae</i>)	HGENLLGQG VKDENVETNKRKREQISTDNEAKMQIQEEKSPKKRKRSSKANK
<i>Sun2NLS</i> (pAK99) (<i>Homo sapiens</i>)	GS SSSSSGSSVAGSQSTLFKDSPLRTLKRKSSNMKRLSPAPQLGPSSDAHTSYYESLVHESWFPPRSSLE ELHGDANWGEDLRVRRRRG
<i>Lem2NLS</i> (pAK100) (<i>Mus musculus</i>)	GS GLSYPPHAGPGPLRRRAS
<i>LBRNLS</i> (pAK102) (<i>Homo sapiens</i>)	GS KPLTSFRQRKGGSTSSPSRRRGSRSRSRSPGRPPKSARRSASASHQADIKEARREV
<i>Lap2θNLS</i> (pAK103) (<i>Rattus norvegicus</i>)	GS RESTRGSRRTPRKRIVETS
TM-GFP (pAK01) (TM: synthetic)	MPRYAQLVSDVRETQPFASIALSKLPREHGCGYQIRQERYWLGFLIPCIISLFIAGNWLWIFLHAIFPKMI DIGRKTKNASRTSGSPGLQEFDIKLIDTVLD MSKGEELFTGVVPILVELDGDVNGHKFSVSGEGEDATY GKLTIKFICTTGKLPVPWPTLVTTFGYGVQCFARYPDHMKQHDFFSKAMPEGYVQERTIFFKDDGNYKT RAEVKFEGDTLVNRIELKGIDFKEDGNILGHKLEYNYNSHNVYIMADKQKNGIKVNFKIRHNIEDGSVQLA DHYQQNTPIGDPVLLPDNHYLSTQSALSKDPNEKRDHMVLLFEVTAAGITHGMDELYK
TM-L-NLS-GFP (TM: synthetic, L: Heh2 ₁₄₅₋₃₁₅) (At “_NLS_” the below indicated NLS sequences are introduced)	MPRYAQLVSDVRETQPFASIALSKLPREHGCGYQIRQERYWLGFLIPCIISLFIAGNWLWIFLHAIFPKMI DIGRKTKNASRTSGSKSDGKATSADLTSELTVEELHKKDSSDDKPRVKELPKPELNLKVSNEFLAQLNKE LASAATENYDHSIKSTDLSSIRIETEEPVGPSTGAETRNESEVMENINLEVQPEVKEAKEELTKISETFDNQD EEDTSRLSSKNIRSPKGRTRHFIANKTKRGIDIMKPFI_NLS_DIKLIDTVLD MSKGEELFTGVVPILVELD GDVNGHKFSVSGEGEDATYGKLTIKFICTTGKLPVPWPTLVTTFGYGVQCFARYPDHMKQHDFFSKA MPEGYVQERTIFFKDDGNYKTRAEVKFEGDTLVNRIELKGIDFKEDGNILGHKLEYNYNSHNVYIMADKQ KNGIKVNFKIRHNIEDGSVQLADHYQQNTPIGDPVLLPDNHYLSTQSALSKDPNEKRDHMVLLFEVTAAG ITHGMDELYK
<i>Pom121NLS</i> ₂₉₀₋₄₈₄ (pAK54/pAK105) (<i>Rattus norvegicus</i>)	ALKEKKKRTVAEEDQLHLDGQENKRRRHDSGSGHSAFEPLVANGVPAAFVPKPGSLKRSLASQSSDDH LNKRSRTSSVSLTSTCTGGIPSSRNAITSSYSSTRGVSQLWKRSGPTSSPFSSPASSRSQTPERPAKKTREE EPCHQSSSAPLVTDKESPGKEVTDPATGKQQLWTSPTPGSSGQQRKKIQ
<i>h2NLS</i> (pAK55) (<i>Saccharomyces cerevisiae</i>)	AVKDENVETNKRKREQISTDNEAKMQIQEEKSPKKRKRSSKANKPPESPPQS
<i>Pom121NLS</i> ₂₉₀₋₃₂₆ (pAK95/pAK96/pAK97) (<i>Rattus norvegicus</i>)	ALKEKKKRTVAEEDQLHLDGQENKRRRHDSGSGHSA
<i>Pom121NLS</i> ₃₂₃₋₄₈₄ (pAK107) (<i>Rattus norvegicus</i>)	AGHSAFEPLVANGVPAAFVPKPGSLKRSLASQSSDDHNLKRSRTSSVSLTSTCTGGIPSSRNAITSSYSST RGVSQLWKRSGPTSSPFSSPASSRSQTPERPAKKTREEEPCHQSSSAPLVTDKESPGKEVTDPATGKQQS LWTSPTPGSSGQQRKKIQ
TM-NLS-L44/53-GFP (At “_NLS_” the below indicated NLS sequences are introduced)	MPRYAQLVSDVRETQPFASIALSKLPREHGCGYQIRQERYWLGFLIPCIISLFIAGNWLWIFLHAIFPKMI DIGRKTKNASRTSGSKSDGKATSRGIDIMKPFI_NLS_DIKLIDTVLD MSKGEELFTGVVPILVELDGDVNG HKFSVSGEGEDATYGKLTIKFICTTGKLPVPWPTLVTTFGYGVQCFARYPDHMKQHDFFSKAMPEGY VQERTIFFKDDGNYKTRAEVKFEGDTLVNRIELKGIDFKEDGNILGHKLEYNYNSHNVYIMADKQKNGIKV NFKIRHNIEDGSVQLADHYQQNTPIGDPVLLPDNHYLSTQSALSKDPNEKRDHMVLLFEVTAAGITHG MDELYK
<i>Pom121NLS</i> ₂₉₀₋₄₈₄ (pAK93) (<i>Rattus norvegicus</i>)	ALKEKKKRTVAEEDQLHLDGQENKRRRHDSGSGHSAFEPLVANGVPAAFVPKPGSLKRSLASQSSDDH NKRRTSSVSLTSTCTGGIPSSRNAITSSYSSTRGVSQLWKRSGPTSSPFSSPASSRSQTPERPAKKTREE PCHQSSSAPLVTDKESPGKEVTDPATGKQQLWTSPTPGSSGQQRKKIQ
<i>h2NLS</i> (pAK94) (<i>Saccharomyces cerevisiae</i>)	AVKDENVETNKRKREQISTDNEAKMQIQEEKSPKKRKRSSKANKPPESPPQS
<i>Pom121NLS</i> ₂₉₀₋₃₂₆ (pAK104)	ALKEKKKRTVAEEDQLHLDGQENKRRRHDSGSGHSA

¹ **Green:** GFP, **italics:** transmembrane region, **black:** ID linker region, **red:** NLS region
with in black and **black/underlined** the P2 and P2' position respectively

Table S7: Amino acid sequences of reporter proteins in HEK293T cells¹

<p>2GFP (pAK111)</p> <p>Between brackets is absent in 2GFP-NLS fusions where the below indicated NLS sequences are introduced</p>	<p>MVSKGEELFTGVVPILVELDGDVNGHKFSVSGEGEGDATYGKLTLFICTTGKLPVPWPTLVTTLTYGVQCFSRYPDHMKQHDFFKSAMPEGYVQERTIFFKDDGNYKTRAEVKFEGDTLVNRIELKGIDFKEDGNILGHKLEYNYNSHNVYIMADKQKNGIKVNFKIRHNIEDGSVQLADHYQQNTPIGDGPVLLPDNHYLSTQSALSKDPNEKRDHMLLEFVTAAGITLGMDELYKSGLSRAMVSKGEELFTGVVPILVELDGDVNGHKFSVSGEGEGDATYGKLTLFICTTGKLPVPWPTLVTTLTYGVQCFSRYPDHMKQHDFFKSAMPEGYVQERTIFFKDDGNYKTRAEVKFEGDTLVNRIELKGIDFKEDGNILGHKLEYNYNSHNVYIMADKQKNGIKVNFKIRHNIEDGSVQLADHYQQNTPIGDGPVLLPDNHYLSTQSALSKDPNEKRDHMLLEFVTAAGITLGMDELYKQASNS(AVDGTAGPGSTGSR)</p>
<p><i>h2NLS</i> (pAK112) <i>(Saccharomyces cerevisiae)</i></p>	<p>VKDENVETN<u>K</u>RKREQISTDNEAKMQIQEEKSPKKRRKKRSSKANKPPESPQQS</p>
<p><i>Pom121NLS</i>₂₉₀₋₄₈₄ (pAK113/pAK114/ pAK115) <i>(Rattus norvegicus)</i></p>	<p>ALKE<u>K</u>KRRTVAEEDQLHLDGQENKRRRHDSGSGHSAFEPLVANGVPAAFVPKPGSLKRSLASQSSDDHLNKRSTSSVSSLTSTCTGGIPSSSRNAITSSYSSTRGVSQLWKRSGPTSSPSSPASSRSQTPERPAKKTREEPCHQSSSSAPLVTDKESPGKEVTDPATGKQQLWTSPTPGSSGQRRKRIQ</p>
<p><i>Pom121NLS</i>₂₉₀₋₃₂₆ (pAK119/pAK120/pAK121/ pAK122)</p>	<p>ALKE<u>K</u>KRRTVAEEDQLHLDGQENKRRRHDSGSGHSA</p>
<p><i>NP</i>NLS (pAK123)</p>	<p>LAAVKRPAATKKAGQAKKKKLDGSTGSR</p>
<p><i>h2NLS-L-TM</i> <i>Saccharomyces cerevisiae</i></p> <p>Heh2₉₃₋₃₇₈ sequence C-terminally fused to GFP or DAM (pAK37/ pAK49)</p> <p>Underlined: absent in L-TM (pA38/ pAK50)</p> <p>Between brackets: absent in <i>h2NLS-L37-TM</i> (pAK108)</p>	<p>VKDENVETN<u>K</u>RKREQISTDNEAKMQIQEEKSPKKRRKKRSSKANKPPESPQQS(KSDGKATSADLTSELETVEELHKDSSDDKPRVKELPKPELPNLKVSNEFLAQLNKELASAATENYDHSIKSTDLSIRIETEPEVGPSTGAETRNESEVMENINLEVQPEVKEAKEELTKISETFDNQDEEDTSRLSSKKNI)RSPKGRTRHFIANKTKRGI DIMKPFIAHLFIWLWNGAIFLSIICPIFLGLWYREQRIQVGVCYCGHEKPLKSLAISAFPQTERVDSVLQAYR</p>

¹ **Green:** GFP, **italics:** transmembrane region, **black:** ID linker region, **red:** NLS region with in black and black/underlined the P2 and P2' position, respectively